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By Michael Lassner

April 14, 2000

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RE: New Patent Application Transmittal

Sir:

Kindly award a filing date and serial number under 35 USC 111 to the patent application based upon the enclosed specification (and any drawings). Declaration and filing fee are deferred. Please direct all correspondence to the undersigned at the address indicated below.

INVENTOR: Michael Lassner  
1920 Fifth Street  
Davis, CA 95616

Citizenship: US

INVENTOR: Beth Savidge  
1920 Fifth Street  
Davis, CA 95616

Citizenship: US

INVENTOR: James D. Weiss  
800 N. Lindbergh Blvd  
St. Louis, MO 63167

Citizenship: US

INVENTOR: Dusty Post-Beitenmiller  
800 N. Lindbergh Blvd  
St. Louis, MO 63167

Citizenship: US

DEPARTMENT OF COMMERCE

TITLE: NUCLEIC ACID SEQUENCES TO PROTEINS INVOLVED IN  
ISOPRENOID SYNTHESIS

[X] Specification (47 total pages)

[X] 26 Sheets of Drawings

[X] This application claims priority to: a US application filed April 15, 1999 as US Serial No. 60/129,899 and US application filed July 30, 1999 as US Serial No. 60/146,461

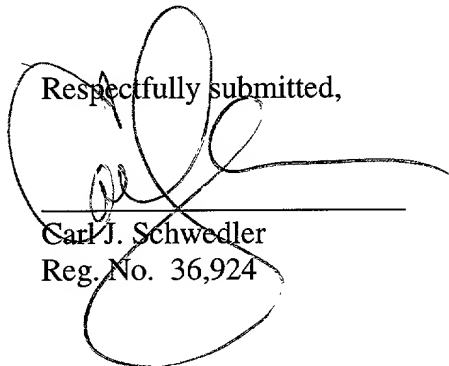
[x ] Sequence Listing (45 pages)

Computer readable form of Sequence Listing as required by 37 CFR 1.821 through 1.825.

It is hereby stated that the content of the paper and computer readable form are the same (§ 1.821(f)).

All correspondence regarding this application should be sent to:

Calgene LLC  
1920 Fifth Street  
Davis, CA 95616  
(530) 753-6313

Respectfully submitted,  
  
Carl J. Schwedler  
Reg. No. 36,924

Enclosure

DEPARTMENT OF COMMERCE

NUCLEIC ACID SEQUENCES TO PROTEINS INVOLVED IN TOCOPHEROL  
SYNTHESIS

5

**INTRODUCTION**

This application claims the benefit of the filing date of the provisional Application U.S. Serial Number 60/129,899, filed April 15, 1999, and the provisional Application, U.S. Serial 10 Number 60/146,461, filed July 30, 1999.

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**TECHNICAL FIELD**

The present invention is directed to nucleic acid and amino acid sequences and constructs, and methods related thereto.

**BACKGROUND**

Isoprenoids are ubiquitous compounds found in all living organisms. Plants synthesize a diverse array of greater than 22,000 isoprenoids (Connolly and Hill (1992) *Dictionary of Terpenoids*, Chapman and Hall, New York, NY). In plants, isoprenoids play essential roles in particular cell functions such as production of sterols, contributing to eukaryotic membrane architecture, acyclic polyprenoids found in the side chain of ubiquinone and plastoquinone, growth regulators like abscisic acid, gibberellins, brassinosteroids or the photosynthetic pigments chlorophylls and carotenoids. Although the physiological role of other plant isoprenoids is less evident, like that of the vast array of secondary metabolites, some are known to play key roles mediating the adaptative responses to different environmental challenges. In spite of the remarkable diversity of structure and function, all isoprenoids originate from a single metabolic precursor, isopentenyl diphosphate (IPP) (Wright, (1961) *Annu. Rev. Biochem.* 20:525-548; and Spurgeon and Porter, (1981) in Biosynthesis of Isoprenoid Compounds., Porter and Spurgeon eds (John Wiley, New York) Vol. 1, pp1-46).

A number of unique and interconnected biochemical pathways derived from the isoprenoid pathway leading to secondary metabolites, including tocopherols, exist in chloroplasts of higher plants. Tocopherols not only perform vital functions in plants, but are also important from mammalian nutritional perspectives. In plastids, tocopherols account for up to 40% of the total quinone pool.

Tocopherols and tocotrienols (unsaturated tocopherol derivatives) are well known antioxidants, and play an important role in protecting cells from free radical damage, and in the prevention of many diseases, including cardiac disease, cancer, cataracts, retinopathy, Alzheimer's disease, and neurodegeneration, and have been shown to have beneficial effects on symptoms of arthritis, and in anti-aging. Vitamin E is used in chicken feed for improving the shelf life, appearance, flavor, and oxidative stability of meat, and to transfer tocots from feed to eggs. Vitamin E has been shown to be essential for normal reproduction, improves overall performance, and enhances immunocompetence in livestock animals. Vitamin E supplement in animal feed also imparts oxidative stability to milk products.

The demand for natural tocopherols as supplements has been steadily growing at a rate of 10-20% for the past three years. At present, the demand exceeds the supply for natural tocopherols, which are known to be more biopotent than racemic mixtures of synthetically produced tocopherols. Naturally occurring tocopherols are all *d*-stereomers, whereas synthetic  $\alpha$ -tocopherol is a mixture of eight *d,l*- $\alpha$ -tocopherol isomers, only one of which (12.5%) is identical to the natural *d*- $\alpha$ -tocopherol. Natural *d*- $\alpha$ -tocopherol has the highest vitamin E activity (1.49 IU/mg) when compared to other natural tocopherols or tocotrienols. The synthetic  $\alpha$ -tocopherol has a vitamin E activity of 1.1 IU/mg. In 1995, the worldwide market for raw refined tocopherols was \$1020 million; synthetic materials comprised 85-88% of the market, the remaining 12-15% being natural materials. The best sources of natural tocopherols and tocotrienols are vegetable oils and grain products. Currently, most of the natural Vitamin E is produced from  $\gamma$ -tocopherol derived from soy oil processing, which is subsequently converted to  $\alpha$ -tocopherol by chemical modification ( $\alpha$ -tocopherol exhibits the greatest biological activity).

Methods of enhancing the levels of tocopherols and tocotrienols in plants, especially levels of the more desirable compounds that can be used directly, without chemical modification, would be useful to the art as such molecules exhibit better functionality and bioavailability.

In addition, methods for the increased production of other isoprenoid derived compounds in a host plant cell is desirable. Furthermore, methods for the production of particular isoprenoid compounds in a host plant cell is also needed.

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## SUMMARY OF THE INVENTION

The present invention is directed to prenyltransferase (PT), and in particular to PT polynucleotides and polypeptides. The polynucleotides and polypeptides of the present invention include those derived from prokaryotic and eukaryotic sources.

Thus, one aspect of the present invention relates to isolated polynucleotide sequences encoding prenyltransferase proteins. In particular, isolated nucleic acid sequences encoding PT proteins from bacterial and plant sources are provided.

Another aspect of the present invention relates to oligonucleotides which include partial or complete PT encoding sequences.

It is also an aspect of the present invention to provide recombinant DNA constructs which can be used for transcription or transcription and translation (expression) of prenyltransferase. In particular, constructs are provided which are capable of transcription or transcription and translation in host cells.

In another aspect of the present invention, methods are provided for production of prenyltransferase in a host cell or progeny thereof. In particular, host cells are transformed or transfected with a DNA construct which can be used for transcription or transcription and translation of prenyltransferase. The recombinant cells which contain prenyltransferase are also part of the present invention.

In a further aspect, the present invention relates to methods of using polynucleotide and polypeptide sequences to modify the tocopherol content of host cells, particularly in host plant cells. Plant cells having such a modified tocopherol content are also contemplated herein.

The modified plants, seeds and oils obtained by the expression of the prenyltransferases are also considered part of the invention.

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## BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 provides an amino acid sequence alignment between ATPT2, ATPT3, ATPT4,

ATPT8, and ATPT12 are performed using ClustalW.

5 Figure 2 provides a schematic picture of the expression construct pCGN10800.

Figure 3 provides a schematic picture of the expression construct pCGN10801.

Figure 4 provides a schematic picture of the expression construct pCGN10803.

Figure 5 provides a schematic picture of the expression construct pCGN10806.

Figure 6 provides a schematic picture of the expression construct pCGN10807.

10 Figure 7 provides a schematic picture of the expression construct pCGN10808.

Figure 8 provides a schematic picture of the expression construct pCGN10809.

Figure 9 provides a schematic picture of the expression construct pCGN10810.

Figure 10 provides a schematic picture of the expression construct pCGN10811.

Figure 11 provides a schematic picture of the expression construct pCGN10812.

Figure 12 provides a schematic picture of the expression construct pCGN10813.

Figure 13 provides a schematic picture of the expression construct pCGN10814.

Figure 14 provides a schematic picture of the expression construct pCGN10815.

Figure 15 provides a schematic picture of the expression construct pCGN10816.

Figure 16 provides a schematic picture of the expression construct pCGN10817.

Figure 17 provides a schematic picture of the expression construct pCGN10819.

Figure 18 provides a schematic picture of the expression construct pCGN10824.

Figure 19 provides a schematic picture of the expression construct pCGN10825.

Figure 20 provides a schematic picture of the expression construct pCGN10826.

Figure 21 provides an amino acid sequence alignment using ClustalW between the

25 *Synechocystis* sequence knockouts.

Figure 22 provides an amino acid sequence of the ATPT2, ATPT3, ATPT4, ATPT8, and ATPT12 protein sequences from *Arabidopsis* and the slr1736, slr0926, slr1899, slr0056, and the slr1518 amino acid sequences from *Synechocystis*.

Figure 23 provides the results of the enzymatic assay from preparations of wild type

30 *Synechocystis* strain 6803, and *Synechocystis* slr1736 knockout.

Figure 24 provides bar graphs of HPLC data obtained from seed extracts of transgenic *Arabidopsis* containing pCGN10822, which provides of the expression of the ATPT2 sequence, in the sense orientation, from the napin promoter. Provided are graphs for alpha, gamma, and delta tocopherols, as well as total tocopherol for 22 transformed lines, as well as a 5 nontransformed (wildtype) control.

Figure 25 provides a bar graph of HPLC analysis of seed extracts from *Arabidopsis* plants transformed with pCGN10803 (35S-ATPT2, in the antisense orientation), pCGN10802 (line 1625, napin ATPT2 in the sense orientation), pCGN10809 (line 1627, 35S-ATPT3 in the sense orientation), a nontransformed (wt) control, and a empty vector transformed control.

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## DETAILED DESCRIPTION OF THE INVENTION

The present invention provides, *inter alia*, compositions and methods for altering (for example, increasing and decreasing) the tocopherol levels and/or modulating their ratios in host cells. In particular, the present invention provides polynucleotides, polypeptides, and methods of use thereof for the modulation of tocopherol content in host plant cells.

The present invention provides polynucleotide and polypeptide sequences involved in the prenylation of straight chain and aromatic compounds. Straight chain prenyl transferases as used herein comprises sequences which encode proteins involved in the prenylation of straight chain compounds, including, but not limited to, geranyl geranyl pyrophosphate and farnesyl pyrophosphate. Aromatic prenyl transferases, as used herein, comprises sequences which encode proteins involved in the prenylation of aromatic compounds, including, but not limited to, menaquinone, ubiquinone, chlorophyll, and homogentisic acid. The prenyl transferase of the 25 present invention preferably prenylates homogentisic acid.

The biosynthesis of  $\alpha$ -tocopherol in higher plants involves condensation of homogentisic acid and phytolpyrophosphate to form 2-methyl-6-phytylbenzoquinol that can, by cyclization and subsequent methylations (Fiedler et al., 1982, *Planta*, 155: 511-515, Soll et al., 1980, *Arch. Biochem. Biophys.* 204: 544-550, Marshall et al., 1985 *Phytochem.*, 24: 1705-1711, all of which 30 are herein incorporated by reference in their entirety), form various tocopherols. The *Arabidopsis*

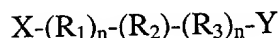
*pds2* mutant identified and characterized by Norris *et al.* (1995), is deficient in tocopherol and plastiquinone-9 accumulation. Further genetic and biochemical analysis suggests that the protein encoded by *PDS2* may be responsible for the prenylation of homogentisic acid. This may be a rate limiting step in tocopherol biosynthesis, and this gene has yet to be isolated. Thus, it is an aspect of the present invention to provide polynucleotides and polypeptides involved in the prenylation of homogentisic acid.

### **Isolated Polynucleotides, Proteins, and Polypeptides**

10 A first aspect of the present invention relates to isolated prenyltransferase polynucleotides. The polynucleotide sequences of the present invention include isolated polynucleotides that encode the polypeptides of the invention having a deduced amino acid sequence selected from the group of sequences set forth in the Sequence Listing and to other polynucleotide sequences closely related to such sequences and variants thereof.

20 The invention provides a polynucleotide sequence identical over its entire length to each coding sequence as set forth in the Sequence Listing. The invention also provides the coding sequence for the mature polypeptide or a fragment thereof, as well as the coding sequence for the mature polypeptide or a fragment thereof in a reading frame with other coding sequences, such as those encoding a leader or secretory sequence, a pre-, pro-, or prepro- protein sequence. The polynucleotide can also include non-coding sequences, including for example, but not limited to, non-coding 5' and 3' sequences, such as the transcribed, untranslated sequences, termination signals, ribosome binding sites, sequences that stabilize mRNA, introns, polyadenylation signals, and additional coding sequence that encodes additional amino acids. For example, a marker sequence can be included to facilitate the purification of the fused polypeptide. Polynucleotides 25 of the present invention also include polynucleotides comprising a structural gene and the naturally associated sequences that control gene expression.

The invention also includes polynucleotides of the formula:



30 wherein, at the 5' end, X is hydrogen, and at the 3' end, Y is hydrogen or a metal, R<sub>1</sub> and R<sub>3</sub> are any nucleic acid residue, n is an integer between 1 and 3000, preferably between 1 and 1000 and

R<sub>2</sub> is a nucleic acid sequence of the invention, particularly a nucleic acid sequence selected from the group set forth in the Sequence Listing and preferably those of SEQ ID NOs: 1, 3, 5, 7, 8, 10, 11, 13-16, 18, 23, 29, 36, and 38. In the formula, R<sub>2</sub> is oriented so that its 5' end residue is at the left, bound to R<sub>1</sub>, and its 3' end residue is at the right, bound to R<sub>3</sub>. Any stretch of nucleic acid residues denoted by either R group, where R is greater than 1, may be either a heteropolymer or a homopolymer, preferably a heteropolymer.

The invention also relates to variants of the polynucleotides described herein that encode for variants of the polypeptides of the invention. Variants that are fragments of the polynucleotides of the invention can be used to synthesize full-length polynucleotides of the invention. Preferred embodiments are polynucleotides encoding polypeptide variants wherein 5 to 10, 1 to 5, 1 to 3, 2, 1 or no amino acid residues of a polypeptide sequence of the invention are substituted, added or deleted, in any combination. Particularly preferred are substitutions, additions, and deletions that are silent such that they do not alter the properties or activities of the polynucleotide or polypeptide.

Further preferred embodiments of the invention that are at least 50%, 60%, or 70% identical over their entire length to a polynucleotide encoding a polypeptide of the invention, and polynucleotides that are complementary to such polynucleotides. More preferable are polynucleotides that comprise a region that is at least 80% identical over its entire length to a polynucleotide encoding a polypeptide of the invention and polynucleotides that are complementary thereto. In this regard, polynucleotides at least 90% identical over their entire length are particularly preferred, those at least 95% identical are especially preferred. Further, those with at least 97% identity are highly preferred and those with at least 98% and 99% identity are particularly highly preferred, with those at least 99% being the most highly preferred.

Preferred embodiments are polynucleotides that encode polypeptides that retain substantially the same biological function or activity as the mature polypeptides encoded by the polynucleotides set forth in the Sequence Listing.

The invention further relates to polynucleotides that hybridize to the above-described sequences. In particular, the invention relates to polynucleotides that hybridize under stringent conditions to the above-described polynucleotides. As used herein, the terms "stringent conditions" and "stringent hybridization conditions" mean that hybridization will generally occur

if there is at least 95% and preferably at least 97% identity between the sequences. An example of stringent hybridization conditions is overnight incubation at 42°C in a solution comprising 50% formamide, 5x SSC (150 mM NaCl, 15 mM trisodium citrate), 50 mM sodium phosphate (pH 7.6), 5x Denhardt's solution, 10% dextran sulfate, and 20 micrograms/milliliter denatured, 5 sheared salmon sperm DNA, followed by washing the hybridization support in 0.1x SSC at approximately 65°C. Other hybridization and wash conditions are well known and are exemplified in Sambrook, *et al.*, Molecular Cloning: A Laboratory Manual, Second Edition, cold Spring Harbor, NY (1989), particularly Chapter 11.

The invention also provides a polynucleotide consisting essentially of a polynucleotide sequence obtainable by screening an appropriate library containing the complete gene for a polynucleotide sequence set forth in the Sequence Listing under stringent hybridization conditions with a probe having the sequence of said polynucleotide sequence or a fragment thereof; and isolating said polynucleotide sequence. Fragments useful for obtaining such a polynucleotide include, for example, probes and primers as described herein.

As discussed herein regarding polynucleotide assays of the invention, for example, polynucleotides of the invention can be used as a hybridization probe for RNA, cDNA, or genomic DNA to isolate full length cDNAs or genomic clones encoding a polypeptide and to isolate cDNA or genomic clones of other genes that have a high sequence similarity to a polynucleotide set forth in the Sequence Listing. Such probes will generally comprise at least 15 bases. Preferably such probes will have at least 30 bases and can have at least 50 bases. Particularly preferred probes will have between 30 bases and 50 bases, inclusive.

The coding region of each gene that comprises or is comprised by a polynucleotide sequence set forth in the Sequence Listing may be isolated by screening using a DNA sequence provided in the Sequence Listing to synthesize an oligonucleotide probe. A labeled oligonucleotide having a sequence complementary to that of a gene of the invention is then used to screen a library of cDNA, genomic DNA or mRNA to identify members of the library which hybridize to the probe. For example, synthetic oligonucleotides are prepared which correspond to the prenyltransferase EST sequences. The oligonucleotides are used as primers in polymerase chain reaction (PCR) techniques to obtain 5' and 3' terminal sequence of prenyl transferase genes. Alternatively, where oligonucleotides of low degeneracy can be prepared from particular

prenyltransferase peptides, such probes may be used directly to screen gene libraries for prenyltransferase gene sequences. In particular, screening of cDNA libraries in phage vectors is useful in such methods due to lower levels of background hybridization.

Typically, a prenyltransferase sequence obtainable from the use of nucleic acid probes  
5 will show 60-70% sequence identity between the target prenyltransferase sequence and the encoding sequence used as a probe. However, lengthy sequences with as little as 50-60% sequence identity may also be obtained. The nucleic acid probes may be a lengthy fragment of the nucleic acid sequence, or may also be a shorter, oligonucleotide probe. When longer nucleic acid fragments are employed as probes (greater than about 100 bp), one may screen at lower  
10 stringencies in order to obtain sequences from the target sample which have 20-50% deviation (i.e., 50-80% sequence homology) from the sequences used as probe. Oligonucleotide probes can be considerably shorter than the entire nucleic acid sequence encoding an prenyltransferase enzyme, but should be at least about 10, preferably at least about 15, and more preferably at least about 20 nucleotides. A higher degree of sequence identity is desired when shorter regions are used as opposed to longer regions. It may thus be desirable to identify regions of highly conserved amino acid sequence to design oligonucleotide probes for detecting and recovering other related prenyltransferase genes. Shorter probes are often particularly useful for polymerase chain reactions (PCR), especially when highly conserved sequences can be identified. (See,  
Gould, *et al.*, PNAS USA (1989) 86:1934-1938.).

20

Another aspect of the present invention relates to prenyltransferase polypeptides. Such polypeptides include isolated polypeptides set forth in the Sequence Listing, as well as polypeptides and fragments thereof, particularly those polypeptides which exhibit prenyltransferase activity and also those polypeptides which have at least 50%, 60% or 70% identity, preferably at least 80% identity, more preferably at least 90% identity, and most preferably at least 95% identity to a polypeptide sequence selected from the group of sequences set forth in the Sequence Listing, and also include portions of such polypeptides, wherein such portion of the polypeptide preferably includes at least 30 amino acids and more preferably includes at least 50 amino acids.

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“Identity”, as is well understood in the art, is a relationship between two or more polypeptide sequences or two or more polynucleotide sequences, as determined by comparing the  
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sequences. In the art, "identity" also means the degree of sequence relatedness between polypeptide or polynucleotide sequences, as determined by the match between strings of such sequences. "Identity" can be readily calculated by known methods including, but not limited to, those described in *Computational Molecular Biology*, Lesk, A.M., ed., Oxford University Press, New York (1988); *Biocomputing: Informatics and Genome Projects*, Smith, D.W., ed., Academic Press, New York, 1993; *Computer Analysis of Sequence Data, Part I*, Griffin, A.M. and Griffin, H.G., eds., Humana Press, New Jersey (1994); *Sequence Analysis in Molecular Biology*, von Heinje, G., Academic Press (1987); *Sequence Analysis Primer*, Gribskov, M. and Devereux, J., eds., Stockton Press, New York (1991); and Carillo, H., and Lipman, D., *SIAM J Applied Math*, 48:1073 (1988). Methods to determine identity are designed to give the largest match between the sequences tested. Moreover, methods to determine identity are codified in publicly available programs. Computer programs which can be used to determine identity between two sequences include, but are not limited to, GCG (Devereux, J., et al., *Nucleic Acids Research* 12(1):387 (1984); suite of five BLAST programs, three designed for nucleotide sequences queries (BLASTN, BLASTX, and TBLASTX) and two designed for protein sequence queries (BLASTP and TBLASTN) (Coulson, *Trends in Biotechnology*, 12: 76-80 (1994); Birren, et al., *Genome Analysis*, 1: 543-559 (1997)). The BLAST X program is publicly available from NCBI and other sources (*BLAST Manual*, Altschul, S., et al., NCBI NLM NIH, Bethesda, MD 20894; Altschul, S., et al., *J. Mol. Biol.*, 215:403-410 (1990)). The well known Smith Waterman algorithm can also be used to determine identity.

Parameters for polypeptide sequence comparison typically include the following:

Algorithm: Needleman and Wunsch, *J. Mol. Biol.* 48:443-453 (1970)

Comparison matrix: BLOSSUM62 from Hentikoff and Hentikoff, *Proc. Natl. Acad. Sci USA* 89:10915-10919 (1992)

25 Gap Penalty: 12

Gap Length Penalty: 4

A program which can be used with these parameters is publicly available as the "gap" program from Genetics Computer Group, Madison Wisconsin. The above parameters along with no penalty for end gap are the default parameters for peptide comparisons.

30 Parameters for polynucleotide sequence comparison include the following:

Algorithm: Needleman and Wunsch, J. Mol. Biol. 48:443-453 (1970)

Comparison matrix: matches = +10; mismatches = 0

Gap Penalty: 50

Gap Length Penalty: 3

5 A program which can be used with these parameters is publicly available as the "gap" program from Genetics Computer Group, Madison Wisconsin. The above parameters are the default parameters for nucleic acid comparisons.

The invention also includes polypeptides of the formula:



10 wherein, at the amino terminus, X is hydrogen, and at the carboxyl terminus, Y is hydrogen or a metal, R<sub>1</sub> and R<sub>3</sub> are any amino acid residue, n is an integer between 1 and 1000, and R<sub>2</sub> is an amino acid sequence of the invention, particularly an amino acid sequence selected from the group set forth in the Sequence Listing and preferably those encoded by the sequences provided in SEQ ID NOS: 2, 4, 6, 9, 12, 17, 19-22, 24-28, 30, 32-35, 37, and 39. In the formula, R<sub>2</sub> is oriented so that its amino terminal residue is at the left, bound to R<sub>1</sub>, and its carboxy terminal residue is at the right, bound to R<sub>3</sub>. Any stretch of amino acid residues denoted by either R group, where R is greater than 1, may be either a heteropolymer or a homopolymer, preferably a heteropolymer.

20 Polypeptides of the present invention include isolated polypeptides encoded by a polynucleotide comprising a sequence selected from the group of a sequence contained in the Sequence Listing set forth herein .

The polypeptides of the present invention can be mature protein or can be part of a fusion protein.

25 Fragments and variants of the polypeptides are also considered to be a part of the invention. A fragment is a variant polypeptide which has an amino acid sequence that is entirely the same as part but not all of the amino acid sequence of the previously described polypeptides. The fragments can be "free-standing" or comprised within a larger polypeptide of which the fragment forms a part or a region, most preferably as a single continuous region. Preferred fragments are biologically active fragments which are those fragments that mediate activities of 30 the polypeptides of the invention, including those with similar activity or improved activity or

with a decreased activity. Also included are those fragments that antigenic or immunogenic in an animal, particularly a human.

Variants of the polypeptide also include polypeptides that vary from the sequences set forth in the Sequence Listing by conservative amino acid substitutions, substitution of a residue 5 by another with like characteristics. In general, such substitutions are among Ala, Val, Leu and Ile; between Ser and Thr; between Asp and Glu; between Asn and Gln; between Lys and Arg; or between Phe and Tyr. Particularly preferred are variants in which 5 to 10; 1 to 5; 1 to 3 or one amino acid(s) are substituted, deleted, or added, in any combination.

Variants that are fragments of the polypeptides of the invention can be used to produce 10 the corresponding full length polypeptide by peptide synthesis. Therefore, these variants can be used as intermediates for producing the full-length polypeptides of the invention.

The polynucleotides and polypeptides of the invention can be used, for example, in the transformation of host cells, such as plant host cells, as further discussed herein.

The invention also provides polynucleotides that encode a polypeptide that is a mature protein plus additional amino or carboxyl-terminal amino acids, or amino acids within the mature polypeptide (for example, when the mature form of the protein has more than one polypeptide chain). Such sequences can, for example, play a role in the processing of a protein from a precursor to a mature form, allow protein transport, shorten or lengthen protein half-life, or facilitate manipulation of the protein in assays or production. It is contemplated that cellular enzymes can be used to remove any additional amino acids from the mature protein.

A precursor protein, having the mature form of the polypeptide fused to one or more prosequences may be an inactive form of the polypeptide. The inactive precursors generally are activated when the prosequences are removed. Some or all of the prosequences may be removed prior to activation. Such precursor protein are generally called proproteins.

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### **Plant Constructs and Methods of Use**

Of particular interest is the use of the nucleotide sequences in recombinant DNA constructs to direct the transcription or transcription and translation (expression) of the 30 prenyltransferase sequences of the present invention in a host plant cell. The expression

constructs generally comprise a promoter functional in a host plant cell operably linked to a nucleic acid sequence encoding a prenyltransferase of the present invention and a transcriptional termination region functional in a host plant cell.

A first nucleic acid sequence is "operably linked" or "operably associated" with a second nucleic acid sequence when the sequences are so arranged that the first nucleic acid sequence affects the function of the second nucleic-acid sequence. Preferably, the two sequences are part of a single contiguous nucleic acid molecule and more preferably are adjacent. For example, a promoter is operably linked to a gene if the promoter regulates or mediates transcription of the gene in a cell.

Those skilled in the art will recognize that there are a number of promoters which are functional in plant cells, and have been described in the literature. Chloroplast and plastid specific promoters, chloroplast or plastid functional promoters, and chloroplast or plastid operable promoters are also envisioned.

One set of plant functional promoters are constitutive promoters such as the CaMV35S or FMV35S promoters that yield high levels of expression in most plant organs. Enhanced or duplicated versions of the CaMV35S and FMV35S promoters are useful in the practice of this invention (Odell, *et al.* (1985) *Nature* 313:810-812; Rogers, U.S. Patent Number 5,378,619). In addition, it may also be preferred to bring about expression of the prenyltransferase gene in specific tissues of the plant, such as leaf, stem, root, tuber, seed, fruit, etc., and the promoter chosen should have the desired tissue and developmental specificity.

Of particular interest is the expression of the nucleic acid sequences of the present invention from transcription initiation regions which are preferentially expressed in a plant seed tissue. Examples of such seed preferential transcription initiation sequences include those sequences derived from sequences encoding plant storage protein genes or from genes involved in fatty acid biosynthesis in oilseeds. Examples of such promoters include the 5' regulatory regions from such genes as napin (Kridl *et al.*, *Seed Sci. Res.* 1:209:219 (1991)), phaseolin, zein, soybean trypsin inhibitor, ACP, stearoyl-ACP desaturase, soybean  $\alpha'$  subunit of  $\beta$ -conglycinin (soy 7s, (Chen *et al.*, *Proc. Natl. Acad. Sci.*, 83:8560-8564 (1986))) and oleosin.

It may be advantageous to direct the localization of proteins conferring prenyltransferase to a particular subcellular compartment, for example, to the mitochondrion, endoplasmic

reticulum, vacuoles, chloroplast or other plastidic compartment. For example, where the genes of interest of the present invention will be targeted to plastids, such as chloroplasts, for expression, the constructs will also employ the use of sequences to direct the gene to the plastid. Such sequences are referred to herein as chloroplast transit peptides (CTP) or plastid transit peptides (PTP). In this manner, where the gene of interest is not directly inserted into the plastid, the expression construct will additionally contain a gene encoding a transit peptide to direct the gene of interest to the plastid. The chloroplast transit peptides may be derived from the gene of interest, or may be derived from a heterologous sequence having a CTP. Such transit peptides are known in the art. See, for example, Von Heijne *et al.* (1991) *Plant Mol. Biol. Rep.* 9:104-126; Clark *et al.* (1989) *J. Biol. Chem.* 264:17544-17550; della-Cioppa *et al.* (1987) *Plant Physiol.* 84:965-968; Romer *et al.* (1993) *Biochem. Biophys. Res Commun.* 196:1414-1421; and, Shah *et al.* (1986) *Science* 233:478-481.

Depending upon the intended use, the constructs may contain the nucleic acid sequence which encodes the entire prenyltransferase protein, or a portion thereof. For example, where antisense inhibition of a given prenyltransferase protein is desired, the entire prenyltransferase sequence is not required. Furthermore, where prenyltransferase sequences used in constructs are intended for use as probes, it may be advantageous to prepare constructs containing only a particular portion of a prenyltransferase encoding sequence, for example a sequence which is discovered to encode a highly conserved prenyltransferase region.

The skilled artisan will recognize that there are various methods for the inhibition of expression of endogenous sequences in a host cell. Such methods include, but are not limited to, antisense suppression (Smith, *et al.* (1988) *Nature* 334:724-726), co-suppression (Napoli, *et al.* (1989) *Plant Cell* 2:279-289), ribozymes (PCT Publication WO 97/10328), and combinations of sense and antisense Waterhouse, *et al.* (1998) *Proc. Natl. Acad. Sci. USA* 95:13959-13964. Methods for the suppression of endogenous sequences in a host cell typically employ the transcription or transcription and translation of at least a portion of the sequence to be suppressed. Such sequences may be homologous to coding as well as non-coding regions of the endogenous sequence.

Regulatory transcript termination regions may be provided in plant expression constructs of this invention as well. Transcript termination regions may be provided by the DNA sequence

encoding the prenyltransferase or a convenient transcription termination region derived from a different gene source, for example, the transcript termination region which is naturally associated with the transcript initiation region. The skilled artisan will recognize that any convenient transcript termination region which is capable of terminating transcription in a plant cell may be employed in the constructs of the present invention.

Alternatively, constructs may be prepared to direct the expression of the prenyltransferase sequences directly from the host plant cell plastid. Such constructs and methods are known in the art and are generally described, for example, in Svab, *et al.* (1990) *Proc. Natl. Acad. Sci. USA* 87:8526-8530 and Svab and Maliga (1993) *Proc. Natl. Acad. Sci. USA* 90:913-917 and in U.S.

10 Patent Number 5,693,507.

The prenyltransferase constructs of the present invention can be used in transformation methods with additional constructs providing for the expression of other nucleic acid sequences encoding proteins involved in the production of tocopherols, or tocopherol precursors such as homogentisic acid and/or phytylpyrophosphate. Nucleic acid sequences encoding proteins involved in the production of homogentisic acid are known in the art, and include but are not limited to, 4-hydroxyphenylpyruvate dioxygenase (HPPD, EC 1.13.11.27) described for example, by Garcia, *et al.* ((1999) *Plant Physiol.* 119(4):1507-1516), mono or bifunctional tyrA (described for example by Xia, *et al.* (1992) *J. Gen Microbiol.* 138:1309-1316, and Hudson, *et al.* (1984) *J. Mol. Biol.* 180:1023-1051), Oxygenase, 4-hydroxyphenylpyruvate di- (9CI), 4-Hydroxyphenylpyruvate dioxygenase; p-Hydroxyphenylpyruvate dioxygenase; p-Hydroxyphenylpyruvate hydroxylase; p-Hydroxyphenylpyruvate oxidase; p-Hydroxyphenylpyruvic acid hydroxylase; p-Hydroxyphenylpyruvic hydroxylase; p-Hydroxyphenylpyruvic oxidase), 4-hydroxyphenylacetate, NAD(P)H:oxygen oxidoreductase (1-hydroxylating); 4-hydroxyphenylacetate 1-monoxygenase, and the like. In addition, constructs for the expression of nucleic acid sequences encoding proteins involved in the production of phytylpyrophosphate can also be employed with the prenyltransferase constructs of the present invention. Nucleic acid sequences encoding proteins involved in the production of phytylpyrophosphate are known in the art, and include, but are not limited to geranylgeranylpyrophosphate synthase (GGPPS), geranylgeranylpyrophosphate reductase (GGH), 1-deoxyxylulose-5-phosphate synthase, 1- deoxy-D-xylolose-5-phosphate

reductoisomerase, 4-diphosphocytidyl-2-C-methylerythritol synthase, isopentyl pyrophosphate isomerase.

The prenyltransferase sequences of the present invention find use in the preparation of transformation constructs having a second expression cassette for the expression of additional sequences involved in tocopherol biosynthesis. Additional tocopherol biosynthesis sequences of interest in the present invention include, but are not limited to gamma-tocopherol methyltransferase (Shintani, *et al.* (1998) *Science* 282(5396):2098-2100), tocopherol cyclase, and tocopherol methyltransferase.

A plant cell, tissue, organ, or plant into which the recombinant DNA constructs containing the expression constructs have been introduced is considered transformed, transfected, or transgenic. A transgenic or transformed cell or plant also includes progeny of the cell or plant and progeny produced from a breeding program employing such a transgenic plant as a parent in a cross and exhibiting an altered phenotype resulting from the presence of a prenyltransferase nucleic acid sequence.

Plant expression or transcription constructs having a prenyltransferase as the DNA sequence of interest for increased or decreased expression thereof may be employed with a wide variety of plant life, particularly, plant life involved in the production of vegetable oils for edible and industrial uses. Particularly preferred plants for use in the methods of the present invention include, but are not limited to: *Acacia*, alfalfa, aneth, apple, apricot, artichoke, arugula, asparagus, avocado, banana, barley, beans, beet, blackberry, blueberry, broccoli, brussels sprouts, cabbage, canola, cantaloupe, carrot, cassava, cauliflower, celery, cherry, chicory, cilantro, citrus, clementines, coffee, corn, cotton, cucumber, Douglas fir, eggplant, endive, escarole, eucalyptus, fennel, figs, garlic, gourd, grape, grapefruit, honey dew, jicama, kiwifruit, lettuce, leeks, lemon, lime, Loblolly pine, mango, melon, mushroom, nectarine, nut, oat, oil palm, oil seed rape, okra, onion, orange, an ornamental plant, papaya, parsley, pea, peach, peanut, pear, pepper, persimmon, pine, pineapple, plantain, plum, pomegranate, poplar, potato, pumpkin, quince, radiata pine, radicchio, radish, raspberry, rice, rye, sorghum, Southern pine, soybean, spinach, squash, strawberry, sugarbeet, sugarcane, sunflower, sweet potato, sweetgum, tangerine, tea, tobacco, tomato, triticale, turf, turnip, a vine, watermelon, wheat, yams, and zucchini.

Most especially preferred are temperate oilseed crops. Temperate oilseed crops of interest include, but are not limited to, rapeseed (Canola and High Erucic Acid varieties), sunflower, safflower, cotton, soybean, peanut, coconut and oil palms, and corn. Depending on the method for introducing the recombinant constructs into the host cell, other DNA sequences may be  
5 required. Importantly, this invention is applicable to dicotyledyons and monocotyledons species alike and will be readily applicable to new and/or improved transformation and regulation techniques.

10 Of particular interest, is the use of prenyltransferase constructs in plants to produce plants or plant parts, including, but not limited to leaves, stems, roots, reproductive, and seed, with a modified content of tocopherols in plant parts having transformed plant cells.

For immunological screening, antibodies to the protein can be prepared by injecting rabbits or mice with the purified protein or portion thereof, such methods of preparing antibodies being well known to those in the art. Either monoclonal or polyclonal antibodies can be produced, although typically polyclonal antibodies are more useful for gene isolation. Western analysis may be conducted to determine that a related protein is present in a crude extract of the desired plant species, as determined by cross-reaction with the antibodies to the encoded proteins. When cross-reactivity is observed, genes encoding the related proteins are isolated by screening expression libraries representing the desired plant species. Expression libraries can be constructed in a variety of commercially available vectors, including lambda gt11, as described in Sambrook, *et al.* (*Molecular Cloning: A Laboratory Manual*, Second Edition (1989) Cold Spring Harbor Laboratory, Cold Spring Harbor, New York).

25 To confirm the activity and specificity of the proteins encoded by the identified nucleic acid sequences as prenyltransferase enzymes, *in vitro* assays are performed in insect cell cultures using baculovirus expression systems. Such baculovirus expression systems are known in the art and are described by Lee, *et al.* U.S. Patent Number 5,348,886, the entirety of which is herein incorporated by reference.

30 In addition, other expression constructs may be prepared to assay for protein activity utilizing different expression systems. Such expression constructs are transformed into yeast or prokaryotic host and assayed for prenyltransferase activity. Such expression systems are known in the art and are readily available through commercial sources.

In addition to the sequences described in the present invention, DNA coding sequences useful in the present invention can be derived from algae, fungi, bacteria, mammalian sources, plants, etc. Homology searches in existing databases using signature sequences corresponding to conserved nucleotide and amino acid sequences of prenyltransferase can be employed to 5 isolate equivalent, related genes from other sources such as plants and microorganisms. Searches in EST databases can also be employed. Furthermore, the use of DNA sequences encoding enzymes functionally enzymatically equivalent to those disclosed herein, wherein such DNA sequences are degenerate equivalents of the nucleic acid sequences disclosed herein in accordance with the degeneracy of the genetic code, is also encompassed by the present 10 invention. Demonstration of the functionality of coding sequences identified by any of these methods can be carried out by complementation of mutants of appropriate organisms, such as *Synechocystis*, *Shewanella*, yeast, *Pseudomonas*, *Rhodobacter*, etc., that lack specific biochemical reactions, or that have been mutated. The sequences of the DNA coding regions can be optimized by gene resynthesis, based on codon usage, for maximum expression in particular hosts.

For the alteration of tocopherol production in a host cell, a second expression construct can be used in accordance with the present invention. For example, the prenyltransferase expression construct can be introduced into a host cell in conjunction with a second expression construct having a nucleotide sequence for a protein involved in tocopherol biosynthesis.

The method of transformation in obtaining such transgenic plants is not critical to the instant invention, and various methods of plant transformation are currently available. Furthermore, as newer methods become available to transform crops, they may also be directly applied hereunder. For example, many plant species naturally susceptible to *Agrobacterium* infection may be successfully transformed via tripartite or binary vector methods of 25 *Agrobacterium* mediated transformation. In many instances, it will be desirable to have the construct bordered on one or both sides by T-DNA, particularly having the left and right borders, more particularly the right border. This is particularly useful when the construct uses *A. tumefaciens* or *A. rhizogenes* as a mode for transformation, although the T-DNA borders may find use with other modes of transformation. In addition, techniques of microinjection, DNA

particle bombardment, and electroporation have been developed which allow for the transformation of various monocot and dicot plant species.

Normally, included with the DNA construct will be a structural gene having the necessary regulatory regions for expression in a host and providing for selection of transformant cells. The 5 gene may provide for resistance to a cytotoxic agent, e.g. antibiotic, heavy metal, toxin, etc., complementation providing prototrophy to an auxotrophic host, viral immunity or the like. Depending upon the number of different host species the expression construct or components thereof are introduced, one or more markers may be employed, where different conditions for selection are used for the different hosts.

10 Where *Agrobacterium* is used for plant cell transformation, a vector may be used which may be introduced into the *Agrobacterium* host for homologous recombination with T-DNA or the Ti- or Ri-plasmid present in the *Agrobacterium* host. The Ti- or Ri-plasmid containing the T-DNA for recombination may be armed (capable of causing gall formation) or disarmed (incapable of causing gall formation), the latter being permissible, so long as the *vir* genes are present in the transformed *Agrobacterium* host. The armed plasmid can give a mixture of normal plant cells and gall.

In some instances where *Agrobacterium* is used as the vehicle for transforming host plant cells, the expression or transcription construct bordered by the T-DNA border region(s) will be inserted into a broad host range vector capable of replication in *E. coli* and *Agrobacterium*, there being broad host range vectors described in the literature. Commonly used is pRK2 or derivatives thereof. See, for example, Ditta, *et al.*, (*Proc. Nat. Acad. Sci., U.S.A.* (1980) 77:7347-7351) and EPA 0 120 515, which are incorporated herein by reference. Alternatively, one may insert the sequences to be expressed in plant cells into a vector containing separate replication sequences, one of which stabilizes the vector in *E. coli*, and the other in 25 *Agrobacterium*. See, for example, McBride, *et al.* (*Plant Mol. Biol.* (1990) 14:269-276), wherein the pRiHRI (Jouanin, *et al.*, *Mol. Gen. Genet.* (1985) 201:370-374) origin of replication is utilized and provides for added stability of the plant expression vectors in host *Agrobacterium* cells.

Included with the expression construct and the T-DNA will be one or more markers, 30 which allow for selection of transformed Agrobacterium and transformed plant cells. A

number of markers have been developed for use with plant cells, such as resistance to chloramphenicol, kanamycin, the aminoglycoside G418, hygromycin, or the like. The particular marker employed is not essential to this invention, one or another marker being preferred depending on the particular host and the manner of construction.

5 For transformation of plant cells using *Agrobacterium*, explants may be combined and incubated with the transformed *Agrobacterium* for sufficient time for transformation, the bacteria killed, and the plant cells cultured in an appropriate selective medium. Once callus forms, shoot formation can be encouraged by employing the appropriate plant hormones in accordance with known methods and the shoots transferred to rooting medium for regeneration of plants. The  
10 plants may then be grown to seed and the seed used to establish repetitive generations and for isolation of vegetable oils.

There are several possible ways to obtain the plant cells of this invention which contain multiple expression constructs. Any means for producing a plant comprising a construct having a DNA sequence encoding the expression construct of the present invention, and at least one other construct having another DNA sequence encoding an enzyme are encompassed by the present invention. For example, the expression construct can be used to transform a plant at the same time as the second construct either by inclusion of both expression constructs in a single transformation vector or by using separate vectors, each of which express desired genes. The second construct can be introduced into a plant which has already been transformed with the prenyltransferase expression construct, or alternatively, transformed plants, one expressing the prenyltransferase construct and one expressing the second construct, can be crossed to bring the constructs together in the same plant.  
20

The nucleic acid sequences of the present invention can be used in constructs to provide for the expression of the sequence in a variety of host cells, both prokaryotic eukaryotic. Host  
25 cells of the present invention preferably include monocotyledenous and dicotyledenous plant cells.

In general, the skilled artisan is familiar with the standard resource materials which describe specific conditions and procedures for the construction, manipulation and isolation of macromolecules (e.g., DNA molecules, plasmids, etc.), generation of recombinant organisms and  
30 the screening and isolating of clones, (see for example, Sambrook *et al.*, *Molecular Cloning: A*

*Laboratory Manual*, Cold Spring Harbor Press (1989); Maliga *et al.*, *Methods in Plant Molecular Biology*, Cold Spring Harbor Press (1995), the entirety of which is herein incorporated by reference; Birren *et al.*, *Genome Analysis: Analyzing DNA*, 1, Cold Spring Harbor, New York, the entirety of which is herein incorporated by reference).

5 Methods for the expression of sequences in insect host cells are known in the art. Baculovirus expression vectors are recombinant insect viruses in which the coding sequence for a chosen foreign gene has been inserted behind a baculovirus promoter in place of the viral gene, e.g., polyhedrin (Smith and Summers, U.S. Pat. No., 4,745,051, the entirety of which is incorporated herein by reference). Baculovirus expression vectors are known in the art, and are described for example in Doerfler, *Curr. Top. Microbiol. Immunol.* 131:51-68 (1968); Luckow and Summers, *Bio/Technology* 6:47-55 (1988a); Miller, *Annual Review of Microbiol.* 42:177-199 (1988); Summers, *Curr. Comm. Molecular Biology*, Cold Spring Harbor Press, Cold Spring Harbor, N.Y. (1988); Summers and Smith, *A Manual of Methods for Baculovirus Vectors and Insect Cell Culture Procedures*, Texas Ag. Exper. Station Bulletin No. 1555 (1988), the entireties of which is herein incorporated by reference)

10 15 20 25 Methods for the expression of a nucleic acid sequence of interest in a fungal host cell are known in the art. The fungal host cell may, for example, be a yeast cell or a filamentous fungal cell. Methods for the expression of DNA sequences of interest in yeast cells are generally described in "Guide to yeast genetics and molecular biology", Guthrie and Fink, eds. Methods in enzymology , Academic Press, Inc. Vol 194 (1991) and Gene expression technology", Goeddel ed, Methods in Enzymology, Academic Press, Inc., Vol 185 (1991).

Mammalian cell lines available as hosts for expression are known in the art and include many immortalized cell lines available from the American Type Culture Collection (ATCC, Manassas, VA), such as HeLa cells, Chinese hamster ovary (CHO) cells, baby hamster kidney (BHK) cells and a number of other cell lines. Suitable promoters for mammalian cells are also known in the art and include, but are not limited to, viral promoters such as that from Simian Virus 40 (SV40) (Fiers *et al.*, *Nature* 273:113 (1978), the entirety of which is herein incorporated by reference), Rous sarcoma virus (RSV), adenovirus (ADV) and bovine papilloma virus (BPV). Mammalian cells may also require terminator sequences and poly-A addition sequences. Enhancer sequences which increase expression may also be included and sequences which

promote amplification of the gene may also be desirable (for example methotrexate resistance genes).

Vectors suitable for replication in mammalian cells are well known in the art, and may include viral replicons, or sequences which insure integration of the appropriate sequences encoding epitopes into the host genome. Plasmid vectors that greatly facilitate the construction of recombinant viruses have been described (see, for example, Mackett *et al*, *J Virol.* 49:857 (1984); Chakrabarti *et al.*, *Mol. Cell. Biol.* 5:3403 (1985); Moss, In: *Gene Transfer Vectors For Mammalian Cells* (Miller and Calos, eds., Cold Spring Harbor Laboratory, N.Y., p. 10, (1987); all of which are herein incorporated by reference in their entirety).

The invention now being generally described, it will be more readily understood by reference to the following examples which are included for purposes of illustration only and are not intended to limit the present invention.

## EXAMPLES

### Example 1: Identification of Prenyltransferase Sequences

PSI-BLAST (Altschul, *et al.* (1997) *Nuc Acid Res* 25:3389-3402) profiles were generated for both the straight chain and aromatic classes of prenyltransferases. To generate the straight chain profile, a prenyl- transferase from *Porphyra purpurea* (Genbank accession 1709766) was used as a query against the NCBI non-redundant protein database. The *E. coli* enzyme involved in the formation of ubiquinone, ubiA (genbank accession 1790473) was used as a starting sequence to generate the aromatic prenyltransferase profile. These profiles were used to search public and proprietary DNA and protein data bases. In *Arabidopsis* seven putative prenyltransferases of the straight-chain class were identified, ATPT1, (SEQ ID NO:9), ATPT7 (SEQ ID NO:10), ATPT8 (SEQ ID NO:11), ATPT9 (SEQ ID NO:13), ATPT10 (SEQ ID NO:14), ATPT11 (SEQ ID NO:15), and ATPT12 (SEQ ID NO:16) and five were identified of the aromatic class, ATPT2 (SEQ ID NO:1), ATPT3 (SEQ ID NO:3), ATPT4 (SEQ ID NO:5), ATPT5 (SEQ ID NO:7), ATPT6 (SEQ ID NO:8). Additional prenyltransferase sequences from

other plants related to the aromatic class of prenyltransferases, such as soy (SEQ ID NOs: 19-23, the deduced amino acid sequence of SEQ ID NO:23 is provided in SEQ ID NO:24) and maize (SEQ ID NOs:25-29, and 31) are also identified. The deduced amino acid sequence of ZMPT5 (SEQ ID NO:29) is provided in SEQ ID NO:30.

5 Searches are performed on a Silicon Graphics Unix computer using additional Bioaccelerator hardware and GenWeb software supplied by Compugen Ltd. This software and hardware enables the use of the Smith-Waterman algorithm in searching DNA and protein databases using profiles as queries. The program used to query protein databases is profilesearch. This is a search where the query is not a single sequence but a profile based on a multiple 10 alignment of amino acid or nucleic acid sequences. The profile is used to query a sequence data set, i.e., a sequence database. The profile contains all the pertinent information for scoring each position in a sequence, in effect replacing the "scoring matrix" used for the standard query searches. The program used to query nucleotide databases with a protein profile is tprofilesearch. Tprofilesearch searches nucleic acid databases using an amino acid profile query. As the search is running, sequences in the database are translated to amino acid sequences in six reading frames. The output file for tprofilesearch is identical to the output file for profilesearch except for an additional column that indicates the frame in which the best alignment occurred.

15  
20  
25

The Smith-Waterman algorithm, (Smith and Waterman (1981) *supra*), is used to search for similarities between one sequence from the query and a group of sequences contained in the database. E score values as well as other sequence information, such as conserved peptide sequences are used to identify related sequences.

25 To obtain the entire coding region corresponding to the *Arabidopsis* prenyltransferase sequences, synthetic oligo-nucleotide primers are designed to amplify the 5' and 3' ends of partial cDNA clones containing prenyltransferase sequences. Primers are designed according to the respective *Arabidopsis* prenyltransferase sequences and used in Rapid Amplification of

cDNA Ends (RACE) reactions (Frohman *et al.* (1988) *Proc. Natl. Acad. Sci. USA* 85:8998-9002) using the Marathon cDNA amplification kit (Clontech Laboratories Inc, Palo Alto, CA).

Additional BLAST searches are performed using the ATPT2 sequence, a sequence in the class of aromatic prenyl transferases. Additional sequences are identified in soybean libraries

that are similar to the ATPT2 sequence. The additional soybean sequence demonstrates 80% identity and 91% similarity at the amino acid sequence.

Amino acid sequence alignments between ATPT2 (SEQ ID NO:2), ATPT3 (SEQ ID NO:4), ATPT4 (SEQ ID NO:6), ATPT8 (SEQ ID NO:12), and ATPT12 (SEQ ID NO:17) are 5 performed using ClustalW (Figure 1), and the percent identity and similarities are provided in Table 1 below.

**Table 1:**

		ATPT2	ATPT3	ATPT4	ATPT8	ATPT12
ATPT2	% Identity		12	13	11	15
	% similar		25	25	22	32
	% Gap		17	20	20	9
ATPT3	% Identity			12	6	22
	% similar			29	16	38
	% Gap			20	24	14
ATPT4	% Identity				9	14
	% similar				18	29
	% Gap				26	19
ATPT8	% Identity					7
	% similar					19
	% Gap					20
ATPT12	% Identity					
	% similar					
	% Gap					

10

**Example 2: Preparation of Expression Constructs**

A plasmid containing the napin cassette derived from pCGN3223 (described in USPN 5,639,790, the entirety of which is incorporated herein by reference) was modified to make it

more useful for cloning large DNA fragments containing multiple restriction sites, and to allow the cloning of multiple napin fusion genes into plant binary transformation vectors. An adapter comprised of the self annealed oligonucleotide of sequence

CGCGATTAAATGGCGGCCCTGCAGGCAGGCCCTGCAGGGCGGCCATTAAAT

5 (SEQ ID NO:40) was ligated into the cloning vector pBC SK+ (Stratagene) after digestion with the restriction endonuclease BssHII to construct vector pCGN7765. Plamids pCGN3223 and pCGN7765 were digested with NotI and ligated together. The resultant vector, pCGN7770, contains the pCGN7765 backbone with the napin seed specific expression cassette from pCGN3223.

10 The cloning cassette, pCGN7787, essentially the same regulatory elements as pCGN7770, with the exception of the napin regulatory regions of pCGN7770 have been replaced with the double CAMV 35S promoter and the tml polyadenylation and transcriptional termination region.

A binary vector for plant transformation, pCGN5139, was constructed from pCGN1558 (McBride and Summerfelt, (1990) Plant Molecular Biology, 14:269-276). The polylinker of pCGN1558 was replaced as a HindIII/Asp718 fragment with a polylinker containing unique restriction endonuclease sites, AscI, PacI, XbaI, SwaI, BamHI, and NotI. The Asp718 and HindIII restriction endonuclease sites are retained in pCGN5139.

20 A series of turbo binary vectors are constructed to allow for the rapid cloning of DNA sequences into binary vectors containing transcriptional initiation regions (promoters) and transcriptional termination regions.

The plasmid pCGN8618 was constructed by ligating oligonucleotides 5'-  
TCGAGGATCCGGCCGCAAGCTCCTGCAGG-3' (SEQ ID NO:41) and 5'-  
TCGACCTGCAGGAAGCTTGGCCGGATCC-3' (SEQ ID NO:42) into SalI/XhoI-  
25 digested pCGN7770. A fragment containing the napin promoter, polylinker and napin 3' region was excised from pCGN8618 by digestion with Asp718I; the fragment was blunt-ended by filling in the 5' overhangs with Klenow fragment then ligated into pCGN5139 that had been digested with Asp718I and HindIII and blunt-ended by filling in the 5' overhangs with Klenow fragment. A plasmid containing the insert oriented so that the napin promoter was closest to the blunted Asp718I site of pCGN5139 and the napin 3' was closest to the blunted HindIII site was subjected

to sequence analysis to confirm both the insert orientation and the integrity of cloning junctions. The resulting plasmid was designated pCGN8622.

The plasmid pCGN8619 was constructed by ligating oligonucleotides 5'-  
TCGACCTGCAGGAAGCTTGC GGCCGCGGATCC -3' (SEQ ID NO:43) and 5'-  
5 TCGAGGATCCGGCCGCAAGCTCCTGCAGG-3' (SEQ ID NO:44) into SalI/XhoI-digested pCGN7770. A fragment containing the napin promoter, polylinker and napin 3' region was removed from pCGN8619 by digestion with Asp718I; the fragment was blunt-ended by filling in the 5' overhangs with Klenow fragment then ligated into pCGN5139 that had been digested with Asp718I and HindIII and blunt-ended by filling in the 5' overhangs with Klenow 10 fragment. A plasmid containing the insert oriented so that the napin promoter was closest to the blunted Asp718I site of pCGN5139 and the napin 3' was closest to the blunted HindIII site was subjected to sequence analysis to confirm both the insert orientation and the integrity of cloning junctions. The resulting plasmid was designated pCGN8623.

The plasmid pCGN8620 was constructed by ligating oligonucleotides 5'-  
TCGAGGATCCGGCCGCAAGCTCCTGCAGGAGCT -3' (SEQ ID NO:45) and 5'-  
CCTGCAGGAAGCTTGC GGCCGCGGATCC-3' (SEQ ID NO:46) into SalI/SacI-digested pCGN7787. A fragment containing the d35S promoter, polylinker and tml 3' region was removed from pCGN8620 by complete digestion with Asp718I and partial digestion with NotI. The fragment was blunt-ended by filling in the 5' overhangs with Klenow fragment then ligated into pCGN5139 that had been digested with Asp718I and HindIII and blunt-ended by filling in the 5' overhangs with Klenow fragment. A plasmid containing the insert oriented so that the d35S promoter was closest to the blunted Asp718I site of pCGN5139 and the tml 3' was closest to the blunted HindIII site was subjected to sequence analysis to confirm both the insert orientation and the integrity of cloning junctions. The resulting plasmid was designated 25 pCGN8624.

The plasmid pCGN8621 was constructed by ligating oligonucleotides 5'-  
TCGACCTGCAGGAAGCTTGC GGCCGCGGATCCAGCT -3' (SEQ ID NO:47) and 5'-  
GGATCCGGCCGCAAGCTCCTGCAGG-3' (SEQ ID NO:48) into SalI/SacI-digested pCGN7787. A fragment containing the d35S promoter, polylinker and tml 3' region was 30 removed from pCGN8621 by complete digestion with Asp718I and partial digestion with NotI.

The fragment was blunt-ended by filling in the 5' overhangs with Klenow fragment then ligated into pCGN5139 that had been digested with Asp718I and HindIII and blunt-ended by filling in the 5' overhangs with Klenow fragment. A plasmid containing the insert oriented so that the d35S promoter was closest to the blunted Asp718I site of pCGN5139 and the tml 3' was closest 5 to the blunted HindIII site was subjected to sequence analysis to confirm both the insert orientation and the integrity of cloning junctions. The resulting plasmid was designated pCGN8625.

The plasmid construct pCGN8640 is a modification of pCGN8624 described above. A 938bp PstI fragment isolated from transposon Tn7 which encodes bacterial spectinomycin and 10 streptomycin resistance (Fling et al. (1985), *Nucleic Acids Research* 13(19):7095-7106), a determinant for *E. coli* and *Agrobacterium* selection, was blunt ended with Pfu polymerase. The blunt ended fragment was ligated into pCGN8624 that had been digested with SpeI and blunt ended with Pfu polymerase. The region containing the PstI fragment was sequenced to confirm both the insert orientation and the integrity of cloning junctions.

The spectinomycin resistance marker was introduced into pCGN8622 and pCGN8623 as follows. A 7.7 Kbp AvrII-SnaBI fragment from pCGN8640 was ligated to a 10.9 Kbp AvrII-SnaBI fragment from pCGN8623 or pCGN8622, described above. The resulting plasmids were pCGN8641 and pCGN8643, respectively.

The plasmid pCGN8644 was constructed by ligating oligonucleotides 5'-  
20 GATCACCTGCAGGAAGCTTGCAGGCCGCGATCCAATGCA-3' (SEQ ID NO:49) and 5'-  
TTGGATCCCGCGCCGCAAGCTTCCTGCAGGT-3' (SEQ ID NO:50) into BamHI-PstI  
digested pCGN8640.

Synthetic oligonucleotides were designed for use in Polymerase Chain Reactions (PCR) to amplify the coding sequences of ATPT2, ATPT3, ATPT4, ATPT8, and ATPT12 for the preparation 25 of expression constructs and are provided in Table 2 below.

**Table 2:**

Name	Restriction Site	Sequence	SEQ ID NO:
ATPT2	5' NotI	GGATCCGGCCGCACAATGGAGTC TCTGCTCTAGTTCT	51
ATPT2	3' SseI	GGATCCTGCAGGTCACTCAAAAAAA GGTAACAGCAAGT	52
ATPT3	5' NotI	GGATCCGGCCGCACAATGGCGTT	53

		TTTTGGGCTCTCCCGTGT	
ATPT3	3' SseI	GGATCCTGCAGGTTATTGAAA CTTCCAAGTACAAC	54
ATPT4	5' NotI	GGATCCGGGCCGCACAATGTGGCG AAGATCTGTGTT	55
ATPT4	3' SseI	GGATCCTGCAGGTACATGGAGAGTAG AAGGAAGGAGCT	56
ATPT8	5' NotI	GGATCCGGGCCGCACAATGGTACT TGCCGAGGTTCCAAAGCTTGCCTCT	57
ATPT8	3' SseI	GGATCCTGCAGGTCACTTGTTC GTGATGACTCTAT	58
ATPT12	5' NotI	GGATCCGGGCCGCACAATGACTTC GATTCTAACACT	59
ATPT12	3' SseI	GGATCCTGCAGGTCAAGTGTGCGAT GCTAATGCCGT	60

The coding sequences of ATPT2, ATPT3, ATPT4, ATPT8, and ATPT12 were all amplified using the respective PCR primers shown in Table 2 above and cloned into the TopoTA vector (Invitrogen). Constructs containing the respective prenyltransferase sequences were digested with NotI and Sse8387I and cloned into the turbobinary vectors described above.

The sequence encoding ATPT2 prenyltransferase was cloned in the sense orientation into pCGN8640 to produce the plant transformation construct pCGN10800 (Figure 2). The ATPT2 sequence is under control of the 35S promoter.

The ATPT2 sequence was also cloned in the antisense orientation into the construct pCGN8641 to create pCGN10801 (Figure 3). This construct provides for the antisense expression of the ATPT2 sequence from the napin promoter.

The ATPT2 coding sequence was also cloned in the antisense orientation into the vector pCGN8643 to create the plant transformation construct pCGN10802

The ATPT2 coding sequence was also cloned in the antisense orientation into the vector pCGN8644 to create the plant transformation construct pCGN10803 (Figure 4).

The ATPT4 coding sequence was cloned into the vector pCGN864 to create the plant transformation construct pCGN10806 (Figure 5). The ATPT2 coding sequence was cloned into the vector pCGN864 to create the plant transformation construct pCGN10807(Figure 6). The ATPT3 coding sequence was cloned into the vector pCGN864 to create the plant transformation construct pCGN10808 (Figure 7). The ATPT3 coding sequence was cloned in the sense orientation into the

vector pCGN8640 to create the plant transformation construct pCGN10809 (Figure 8). The ATPT3 coding sequence was cloned in the antisense orientation into the vector pCGN8641 to create the plant transformation construct pCGN10810 (Figure 9). The ATPT3 coding sequence was cloned into the vector pCGN8643 to create the plant transformation construct pCGN10811 (Figure 10). The 5 ATPT3 coding sequence was cloned into the vector pCGN8640 to create the plant transformation construct pCGN10812 (Figure 11). The ATPT4 coding sequence was cloned into the vector pCGN8640 to create the plant transformation construct pCGN10813 (Figure 12). The ATPT4 coding sequence was cloned into the vector pCGN8643 to create the plant transformation construct pCGN10814 (Figure 13). The ATPT4 coding sequence was cloned into the vector pCGN8641 to 10 create the plant transformation construct pCGN10815 (Figure 14). The ATPT4 coding sequence was cloned in the antisense orientation into the vector pCGN8644 to create the plant transformation construct pCGN10816 (Figure 15). The ATPT2 coding sequence was cloned into the vector pCGN???? to create the plant transformation construct pCGN10817 (Figure 16). The ATPT8 coding sequence was cloned in the sense orientation into the vector pCGN8643 to create the plant transformation construct pCGN10819 (Figure 17). The ATPT12 coding sequence was cloned into the vector pCGN8644 to create the plant transformation construct pCGN10824 (Figure 18). The ATPT12 coding sequence was cloned into the vector pCGN8641 to create the plant transformation construct pCGN10825 (Figure 19). The ATPT8 coding sequence was cloned into the vector pCGN8644 to create the plant transformation construct pCGN10826 (Figure 20).

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### Example 3: Plant Transformation

Transgenic *Brassica* plants are obtained by *Agrobacterium*-mediated transformation as 25 described by Radke *et al.* (*Theor. Appl. Genet.* (1988) 75:685-694; *Plant Cell Reports* (1992) 11:499-505). Transgenic *Arabidopsis thaliana* plants may be obtained by *Agrobacterium*-mediated transformation as described by Valverkens *et al.*, (*Proc. Nat. Acad. Sci.* (1988) 85:5536-5540), or as described by Bent *et al.* ((1994), *Science* 265:1856-1860), or Bechtold *et al.* ((1993), *C.R.Acad.Sci, Life Sciences* 316:1194-1199). Other plant species may be similarly 30 transformed using related techniques.

Alternatively, microprojectile bombardment methods, such as described by Klein *et al.* (*BioTechnology* 10:286-291) may also be used to obtain nuclear transformed plants.

5   **Example 4:** Identification of Additional Prenyltransferases

A PSI-Blast profile generated using the *E. coli* ubiA (genbank accession 1790473) sequence was used to analyze the *Synechocystis* genome. This analysis identified 5 open reading frames (ORFs) in the *Synechocystis* genome that were potentially prenyltransferases; slr0926  
10 (annotated as ubiA (4-hydroxybenzoate-octaprenyl transferase, SEQ ID NO:32), sll1899 (annotated as ctaB (cytchrome c oxidase folding protein, SEQ ID NO:33), slr0056 (annotated as g4 (chlorophyll synthase 33 kd subunit, SEQ ID NO:34), slr1518 (annotated as menA (menaquinone biosynthesis protein, SEQ ID NO:35), and slr1736 (annotated as a hypothetical protein of unknown function (SEQ ID NO:36).

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To determine the functionality of these ORFs and their involvement, if any, in the biosynthesis of Tocopherols, knockouts constructs were made to disrupt the ORF identified in *Synechocystis*.

Synthetic oligos were designed to amplify regions from the 5' (5'-  
TAATGTGTACATTGTCGGCCTC (17365') (SEQ ID NO:61) and 5'-  
20 GCAATGTAACATCAGAGATTTGAGACACAACGTGGCTTCCACAATTCCCCGCACC  
GTC (1736kanpr1)) (SEQ ID NO:62) and 3' (5'-AGGCTAATAAGCACAAATGGGA (17363')  
(SEQ ID NO:63) and 5'-GGTATGAGTCAGCAACACCTTCTTCACGAGGCAGACCTCAGC  
GGAATTGGTTAGGTTATCCC (1736kanpr2)) (SEQ ID NO:64) ends of the slr1736 ORF.  
The 1736kanpr1 and 1736kanpr2 oligos contained 20 bp of homology to the slr1736 ORF with  
25 an additional 40 bp of sequence homology to the ends of the kanamycin resistance cassette.  
Separate PCR steps were completed with these oligos and the products were gel purified and  
combined with the kanamycin resistance gene from puc4K (Pharmacia) that had been digested  
with *Hinc*II and gel purified away from the vector backbone. The combined fragments were  
allowed to assemble without oligos under the following conditions: 94°C for 1 min, 55°C for 1  
30 min, 72°C for 1 min plus 5 seconds per cycle for 40 cycles using pfu polymerase in 100ul

reaction volume (Zhao, H and Arnold (1997) *Nucleic Acids Res.* 25(6):1307-1308). One microliter or five microliters of this assembly reaction was then amplified using 5' and 3' oligos nested within the ends of the ORF fragment, so that the resulting product contained 100-200 bp of the 5' end of the *Synechocystis* gene to be knocked out, the kanamycin resistance cassette, and 5 100-200 bp of the 3' end of the gene to be knocked out. This PCR product was then cloned into the vector pGemT easy (Promega) to create the construct pMON21681 and used for *Synechocystis* transformation.

Primers were also synthesized for the preparation of *Synechocystis* knockout constructs for the other sequences using the same method as described above, with the following primers.  
10 The ubiA 5' sequence was amplified using the primers 5'- GGATCCATGGTT  
GCCAAACCCCATC (SEQ ID NO:65) and 5' - GCAATGTAACATCAGAGA  
TTTGAGACACAACG TGGCTTGGGTAAAGCAACAATGACCGGC (SEQ ID NO:66).  
The 3' region was amplified using the synthetic oligonucleotide primers 5'-  
GAATTCTCAAAGCCAGCCCAGTAAC (SEQ ID NO:67) and 5'-GGTATGAGTC  
AGCAACACCTTCTTCACGAGGCAGACCTCAGCGGGTGCAGAAAAGGGTTTCCC (SEQ  
ID NO:68). The amplification products were combined with the kanamycin resistance gene from puc4K (Pharmacia) that had been digested with *Hinc*II and gel purified away from the vector backbone. The annealed fragment was amplified using 5' and 3' oligos nested within the ends of the ORF fragment (5' - CCAGTGGTTAGGCTGTGTGGTC (SEQ ID NO:69) and 5'-  
CTGAGTTGGATGTATTGGATC (SEQ ID NO:70)), so that the resulting product contained 100-200 bp of the 5' end of the *Synechocystis* gene to be knocked out, the kanamycin resistance cassette, and 100-200 bp of the 3' end of the gene to be knocked out. This PCR product was then cloned into the vector pGemT easy (Promega) to create the construct pMON21682 and used for *Synechocystis* transformation.

25 Primers were also synthesized for the preparation of *Synechocystis* knockout constructs for the other sequences using the same method as described above, with the following primers. The s111899 5' sequence was amplified using the primers 5' - GGATCCATGGTTACTT  
CGACAAAAATCC (SEQ ID NO:71) and 5' - GCAATGTAACATCAGAG  
ATTTGAGACACAACGTGGCTTGCTAGGCAACCGCTTAGTAC (SEQ ID NO:72). The 30 3' region was amplified using the synthetic oligonucleotide primers 5'-

GAATTCTAACCCAACAGTAAAGTTCCC (SEQ ID NO:73) and 5'- GGTATGAGTCAGC  
AACACCTTCTCACGAGGCAGACCTCAGCGCCGGCATTGTCTTTACATG (SEQ ID  
NO:74). The amplification products were combined with the kanamycin resistance gene from  
5 puc4K (Pharmacia) that had been digested with *Hinc*II and gel purified away from the vector  
backbone. The annealed fragment was amplified using 5' and 3' oligos nested within the ends of  
the ORF fragment (5'- GGAACCCTTGCAGCCGCTTC (SEQ ID NO:75)  
and 5'- GTATGCCCAACTGGTGCAGAGG (SEQ ID NO:76)), so that the resulting product  
contained 100-200 bp of the 5' end of the *Synechocystis* gene to be knocked out, the kanamycin  
resistance cassette, and 100-200 bp of the 3' end of the gene to be knocked out. This PCR  
10 product was then cloned into the vector pGemT easy (Promega) to create the construct  
pMON21679 and used for *Synechocystis* transformation.

Primers were also synthesized for the preparation of *Synechocystis* knockout constructs  
for the other sequences using the same method as described above, with the following primers.  
The slr0056 5' sequence was amplified using the primers 5'-  
GGATCCATGTCTGACACACAAAATACCG (SEQ ID NO:77) and 5'-  
GCAATGTAACATCAGAGATTTGAGACACAACGTGGCTTCGCCAATACCAGCCACC  
AACAG (SEQ ID NO:78). The 3' region was amplified using the synthetic oligonucleotide  
primers 5'- GAATTCTCAAAT CCCGCATGGCCTAG (SEQ ID NO:79) and 5'-  
GGTATGAGTCAGAACACCTTCTCACGAGGCAGACCTCAGCGGCCTACGGCTTGGA  
CGTGTGGG (SEQ ID NO:80). The amplification products were combined with the kanamycin  
resistance gene from puc4K (Pharmacia) that had been digested with *Hinc*II and gel purified  
away from the vector backbone. The annealed fragment was amplified using 5' and 3' oligos  
nested within the ends of the ORF fragment (5'- CACTTGGATTCCCCTGATCTG (SEQ ID  
NO:81) and 5'- GCAATAACCGCTTGGAAAACG (SEQ ID NO:82)), so that the resulting  
25 product contained 100-200 bp of the 5' end of the *Synechocystis* gene to be knocked out, the  
kanamycin resistance cassette, and 100-200 bp of the 3' end of the gene to be knocked out. This  
PCR product was then cloned into the vector pGemT easy (Promega) to create the construct  
pMON21677 and used for *Synechocystis* transformation.

Primers were also synthesized for the preparation of *Synechocystis* knockout constructs  
30 for the other sequences using the same method as described above, with the following primers.

The slr1518 5' sequence was amplified using the primers 5'- GGATCCATGACCGAAT CTTCGCCCCTAGC (SEQ ID NO:83) and 5'-GCAATGTAACATCAGAGATTGAGACACAACGTGGC TTTCAATCCTAGGTAGCCGAGGCG (SEQ ID NO:84). The 3' region was amplified using the synthetic oligonucleotide primers 5'- GAATTCTTAGCCCAGGCC  
5 AGCCCAGCC (SEQ ID NO:85) and 5'- GGTATGAGTCAGAACACCTTCTCACGA GGCAGACCTCAGCGGGGAATTGATTGTTAATTACC (SEQ ID NO:86). The amplification products were combined with the kanamycin resistance gene from puc4K (Pharmacia) that had been digested with *Hinc*II and gel purified away from the vector backbone. The annealed fragment was amplified using 5' and 3' oligos nested within the ends of the ORF  
10 fragment (5'- GCGATCGCCATTATCGCTTGG (SEQ ID NO:87) and 5'- GCAGACTGGCAATTATCAGTAACG (SEQ ID NO:88)), so that the resulting product contained 100-200 bp of the 5' end of the *Synechocystis* gene to be knocked out, the kanamycin resistance cassette, and 100-200 bp of the 3' end of the gene to be knocked out. This PCR product was then cloned into the vector pGemT easy (Promega) to create the construct pMON21680 and used for *Synechocystis* transformation.

B. Transformation of *Synechocystis*

Cells of *Synechocystis* 6803 were grown to a density of approximately  $2 \times 10^8$  cells per ml and harvested by centrifugation. The cell pellet was re-suspended in fresh BG-11 medium (ATCC Medium 616) at a density of  $1 \times 10^9$  cells per ml and used immediately for transformation. One-hundred microliters of these cells were mixed with 5 ul of mini prep DNA and incubated with light at 30C for 4 hours. This mixture was then plated onto nylon filters resting on BG-11 agar supplemented with TES pH8 and allowed to grow for 12-18 hours. The filters were then transferred to BG-11 agar + TES + 5ug/ml kanamycin and allowed to grow until colonies  
25 appeared within 7-10 days (Packer and Glazer, 1988). Colonies were then picked into BG-11 liquid media containing 5 ug/ml kanamycin and allowed to grow for 5 days. These cells were then transferred to Bg-11 media containing 10ug/ml kanamycin and allowed to grow for 5 days and then transferred to Bg-11 + kanamycin at 25ug/ml and allowed to grow for 5 days. Cells  
30 were then harvested for PCR analysis to determine the presence of a disrupted ORF and also for HPLC analysis to determine if the disruption had any effect on tocopherol levels.

PCR analysis of the *Synechocystis* isolates for slr1736 and sll1899 showed complete segregation of the mutant genome, meaning no copies of the wild type genome could be detected in these strains. This suggests that function of the native gene is not essential for cell function. HPLC analysis of these same isolates showed that the sll1899 strain had no detectable reduction 5 in tocopherol levels. However, the strain carrying the knockout for slr1736 produced no detectable levels of tocopherol.

The amino acid sequences for the *Synechocystis* knockouts are compared using ClustalW, and are provided in Table 3 below. Provided are the percent identities, percent similarity, and the percent gap. The alignment of the sequences is provided in Figure 21.

10

**Table 3:**

	Slr1736	slr0926	sll1899	slr0056	slr1518
slr1736 %identity		14	12	18	11
%similar		29	30	34	26
%gap		8	7	10	5
slr0926 %identity			20	19	14
%similar			39	32	28
%gap			7	9	4
sll1899 %identity				17	13
%similar				29	29
%gap				12	9
slr0056 %identity					15
%similar					31
%gap					8
slr1518 %identity					
%similar					
%gap					

Amino acid sequence comparisons are performed using various *Arabidopsis* prenyltransferase sequences and the *Synechocystis* sequences. The comparisons are presented in

Table 4 below. Provided are the percent identities, percent similarity, and the percent gap. The alignment of the sequences is provided in Figure 22.

**Table 4:**

	ATPT2	slr1736	ATPT3	slr0926	ATPT4	sll1899	ATPT12	slr0056	ATPT8	slr1518
ATPT2	29	9	9	8	8	12	9	7	9	
	46	23	21	20	20	28	23	21	20	
	27	13	28	23	29	11	24	25	24	
	9	13	8	12	13	13	15	8	10	
	19	28	19	28	26	26	33	21	26	
	34	12	34	15	26	10	10	12	10	
ATPT3		23	11	14	13	10	5	11		
		36	26	26	26	21	14	22		
		29	21	31	16	30	30	30		
			12	20	17	20	11	14		
				24	28	33	24	29		
				33	12	10	11	9		
ATPT4				18	11	8	6	7		
				33	23	18	16	19		
				28	19	32	32	33		
					13	17	10	12		
					24	30	23	26		
					27	13	10	11		
ATPT1						52	8	11		
						66	19	26		
						18	25	23		
						9	13			
						23	32			
						10	8			
slr0056										

ATPT8	7
	23
	7
slr1518	

#### 4B. Preparation of the slr1737 Knockout

The *Synechocystis* sp. 6803 slr1737 knockout was constructed by the following method.

The GPS™-1 Genome Priming System (New England Biolabs) was used to insert, by a Tn7 Transposase system, a Kanamycin resistance cassette into *slr1737*. A plasmid from a *Synechocystis* genomic library clone containing 652 base pairs of the targeted orf (*Synechocystis* genome base pairs 1324051 – 1324703; the predicted orf base pairs 1323672 – 1324763, as annotated by Cyanobase) was used as target DNA. The reaction was performed according to the manufacturers protocol. The reaction mixture was then transformed into *E. coli* DH10B electrocompetant cells and plated. Colonies from this transformation were then screened for transposon insertions into the target sequence by amplifying with M13 Forward and Reverse Universal primers, yielding a product of 652 base pairs plus ~1700 base pairs, the size of the transposon kanamycin cassette, for a total fragment size of ~2300 base pairs. After this determination, it was then necessary to determine the approximate location of the insertion within the targeted orf, as 100 base pairs of orf sequence was estimated as necessary for efficient homologous recombination in *Synechocystis*. This was accomplished through amplification reactions using either of the primers to the ends of the transposon, Primer S (5' end) or N (3' end), in combination with either a M13 Forward or Reverse primer. That is, four different primer combinations were used to map each potential knockout construct: Primer S – M13 Forward, Primer S – M13 Reverse, Primer N – M13 Forward, Primer N – M13 Reverse. The construct used to transform *Synechocystis* and knockout slr1737 was determined to consist of a approximately 150 base pairs of slr1737 sequence on the 5' side of the transposon insertion and

approximately 500 base pairs on the 3' side, with the transcription of the orf and kanamycin cassette in the same direction. The nucleic acid sequence of slr1737 is provided in SEQ ID NO:38 the deduced amino acid sequence is provided in SEQ ID NO:39.

Cells of *Synechocystis* 6803 were grown to a density of ~ $2 \times 10^8$  cells per ml and harvested by centrifugation. The cell pellet was re-suspended in fresh BG-11 medium at a density of  $1 \times 10^9$  cells per ml and used immediately for transformation. 100 ul of these cells were mixed with 5 ul of mini prep DNA and incubated with light at 30C for 4 hours. This mixture was then plated onto nylon filters resting on BG-11 agar supplemented with TES ph8 and allowed to grow for 12-18 hours. The filters were then transferred to BG-11 agar + TES + 5ug/ml kanamycin and allowed to grow until colonies appeared within 7-10 days (Packer and Glazer, 1988). Colonies were then picked into BG-11 liquid media containing 5 ug/ml kanamycin and allowed to grow for 5 days. These cells were then transferred to Bg-11 media containing 10ug/ml kanamycin and allowed to grow for 5 days and then transferred to Bg-11 + kanamycin at 25ug/ml and allowed to grow for 5 days. Cells were then harvested for PCR analysis to determine the presence of a disrupted ORF and also for HPLC analysis to determine if the disruption had any effect on tocopherol levels.

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PCR analysis of the *Synechocystis* isolates, using primers to the ends of the *slr1737* orf , showed complete segregation of the mutant genome, meaning no copies of the wild type genome could be detected in these strains. This suggests that function of the native gene is not essential for cell function. HPLC analysis of the strain carrying the knockout for *slr1737* produced no detectable levels of tocopherol.

#### 4C. Phytyl Prenyltransferase Enzyme Assays

[<sup>3</sup>H] Homogentisic acid in 0.1% H<sub>3</sub>PO<sub>4</sub> (specific radioactivity 40 Ci/mmol). Phytyl pyrophosphate was synthesized as described by Joo, *et al.* (1973) *Can J. Biochem.* 51:1527. 2-methyl-6-phytylquinol and 2,3-dimethyl-5-phytylquinol were synthesized as described by Soll, *et al.* (1980) *Phytochemistry* 19:215. Homogentisic acid, α, β, δ, and γ-tocopherol, and tocol, were purchased commercially.

The wild-type strain of *Synechocystis* sp. PCC 6803 was grown in BG11 medium with bubbling air at 30°C under 50 μE.m<sup>-2</sup>.s<sup>-1</sup> fluorescent light, and 70% relative humidity. The growth

medium of slr1736 knock-out (potential PPT) strain of this organism was supplemented with 25  $\mu\text{g mL}^{-1}$  kanamycin. Cells were collected from 0.25 to 1 liter culture by centrifugation at 5000g for 10 min and stored at -80°C.

Total membranes were isolated according to Zak's procedures with some modifications (Zak,  
5 et al. (1999) *Eur J. Biochem* 261:311). Cells were broken on a French press. Before the French  
press treatment, the cells were incubated for 1 hour with lysozyme (0.5%, w/v) at 30 °C in a  
medium containing 7 mM EDTA, 5 mM NaCl and 10 mM Hepes-NaOH, pH 7.4. The  
spheroplasts were collected by centrifugation at 5000 g for 10 min and resuspended at 0.1 - 0.5 mg  
chlorophyll·mL<sup>-1</sup> in 20 mM potassium phosphate buffer, pH 7.8. Proper amount of protease  
10 inhibitor cocktail and DNAase I from Boehringer Mannheim were added to the solution. French  
press treatments were performed two to three times at 100 MPa. After breakage, the cell  
suspension was centrifuged for 10 min at 5000g to pellet unbroken cells, and this was followed by  
centrifugation at 100 000 g for 1 hour to collect total membranes. The final pellet was resuspended  
in a buffer containing 50 mM Tris-HCL and 4 mM MgCl<sub>2</sub>.

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Chloroplast pellets were isolated from 250 g of spinach leaves obtained from local markets.  
Devined leaf sections were cut into grinding buffer (2 1 /250 g leaves) containing 2 mM EDTA, 1  
mM MgCl<sub>2</sub>, 1 mM MnCl<sub>2</sub>, 0.33 M sorbitol, 0.1% ascorbic acid, and 50 mM Hepes at pH 7.5. The  
leaves were homogenized for 3 sec three times in a 1-L blender, and filtered through 4 layers of  
mirocloth. The supernatant was then centrifuged at 5000g for 6 min. The chloroplast pellets were  
resuspended in small amount of grinding buffer (Douce,et al Methods in Chloroplast Molecular  
Biology, 239 (1982)

Chloroplasts in pellets can be broken in three ways. Chloroplast pellets were first aliquoted  
in 1 mg of chlorophyll per tube, centrifuged at 6000 rpm for 2 min in microcentrifuge, and  
grinding buffer was removed. Two hundred microliters of Triton X-100 buffer (0.1% Triton X-  
25 100, 50 mM Tris-HCl pH 7.6 and 4 mM MgCl<sub>2</sub>) or swelling buffer (10 mM Tris pH 7.6 and 4  
mM MgCl<sub>2</sub>) was added to each tube and incubated for ½ hour at 4°C. Then the broken  
chloroplast pellets were used for the assay immediately. In addition, broken chloroplasts can also  
be obtained by freezing in liquid nitrogen and stored at -80°C for ½ hour, then used for the assay.

In some cases chloroplast pellets were further purified with 40%/ 80% percoll gradient to  
30 obtain intact chloroplasts. The intact chloroplasts were broken with swelling buffer, then either

used for assay or further purified for envelope membranes with 20.5% / 31.8% sucrose density gradient (Sol, et al (1980) *supra*). The membrane fractions were centrifuged at 100 000g for 40 min and resuspended in 50 mM Tris-HCl pH 7.6, 4 mM MgCl<sub>2</sub>.

Various amounts of [<sup>3</sup>H]HGA, 40 to 60 µM unlabelled HGA with specific activity in the 5 range of 0.16 to 4 Ci/mmol were mixed with a proper amount of 1M Tris-NaOH pH 10 to adjust pH to 7.6. HGA was reduced for 4 min with a trace amount of solid NaBH<sub>4</sub>. In addition to HGA, standard incubation mixture (final vol 1 mL) contained 50 mM Tris-HCl, pH 7.6, 3-5 mM MgCl<sub>2</sub>, and 100 µM phytyl pyrophosphate. The reaction was initiated by addition of *Synechocystis* total membranes, spinach chloroplast pellets, spinach broken chloroplasts, or spinach envelope 10 membranes. The enzyme reaction was carried out for 2 hour at 23°C or 30°C in the dark or light. The reaction is stopped by freezing with liquid nitrogen, and stored at -80°C or directly by extraction.

A constant amount of tocol was added to each assay mixture and reaction products were extracted with a 2 mL mixture of chloroform/methanol (1:2, v/v) to give a monophasic solution. NaCl solution (2 mL; 0.9%) was added with vigorous shaking. This extraction procedure was repeated three times. The organic layer containing the prenylquinones was filtered through a 20 µ filter, evaporated under N<sub>2</sub>, and then resuspended in 100 µL of ethanol.

The samples were mainly analyzed by Normal-Phase HPLC method (Isocratic 90% Hexane and 10% Methyl-t-butyl ether), and use a Zorbax silica column, 4.6 x 250 mm. The samples were 20 also analyzed by Reversed-Phase HPLC method (Isocratic 0.1% H<sub>3</sub>PO<sub>4</sub> in MeOH), and use a Vydac 201HS54 C18 column; 4.6 x 250 mm coupled with an All-tech C18 guard column. The amount of products were calculated based on the substrate specific radioactivity, and adjusted according to the % recovery based on the amount of internal standard.

The amount of chlorophyll was determined as described in Arnon (1949) *Plant Physiol.* 24:1. 25 Amount of protein was determined by the Bradford method using gamma globulin as a standard (Bradford, (1976) *Anal. Biochem.* 72:248)

Results of the assay demonstrate that 2-Methyl-6-Phytylplastoquinone is produced in the *Synechocystis* slr1736 knockout preparations. The results of the phytyl prenyltransferase enzyme activity assay for the slr1736 knock out are presented in Figure 23.

4D. Complementation of the slr1736 knockout with ATPT2

In order to determine whether ATPT2 could complement the knockout of slr1736 in *Synechocystis 6803* a plasmid was constructed to express the ATPT2 sequence from the TAC promoter. A vector, plasmid psl1211, was obtained from the lab of Dr. Himadri Pakrasi of Washington University, and is based on the plasmid RSF1010 which is a broad host range plasmid (Ng W.-O., Zentella R., Wang, Y., Taylor J-S. A., Pakrasi, H.B. 2000. *phrA*, the major photoreactivating factor in the cyanobacterium *Synechocystis* sp. strain PCC 6803 codes for a cyclobutane pyrimidine dimer specific DNA photolyase. *Arch. Microbiol.* (in press)). The ATPT2 gene was isolated from the vector pCGN10817 by PCR using the following primers.

ATPT2nco.pr 5'-CCATGGATTGAGTAAAGTTGTCGC (SEQ ID NO:89); ATPT2ri.pr- 5'-GAATTCACCTCAAAAAAGGTAACAG (SEQ ID NO:90). These primers will remove approximately 112 BP from the 5' end of the ATPT2 sequence, which is thought to be the chloroplast transit peptide. These primers will also add an NcoI site at the 5' end and an EcoRI site at the 3' end which can be used for sub-cloning into subsequent vectors. The PCR product from using these primers and pCGN10817 was ligated into pGEM T easy and the resulting vector pMON21689 was confirmed by sequencing using the m13forward and m13reverse primers. The NcoI/EcoRI fragment from pMON21689 was then ligated with the EagI/EcoRI and EagI/NcoI fragments from psl1211 resulting in pMON21690. The plasmid pMON21690 was introduced into the slr1736 *Synechocystis 6803* KO strain via conjugation. Cells of sl906 (a helper strain) and DH10B cells containing pMON21690 were grown to log phase (O.D. 600= 0.4) and 1 ml was harvested by centrifugation. The cell pellets were washed twice with a sterile BG-11 solution and resuspended in 200 ul of BG-11. The following was mixed in a sterile eppendorf tube: 50 ul SL906, 50 ul DH10B cells containing pMON21690, and 100 ul of a fresh culture of the slr1736 *Synechocystis 6803* KO strain (O.D. 730 = 0.2-0.4). The cell mixture was immediately transferred to a nitrocellulose filter resting on BG-11 and incubated for 24 hours at 30C and 2500 LUX(50 ue) of light. The filter was then transferred to BG-11 supplemented with 10ug/ml Gentamycin and incubated as above for ~5 days. When colonies appeared, they were picked and grown up in liquid BG-11 + Gentamycin 10 ug/ml. (Elhai, J. and Wolk, P. 1988. Conjugal transfer of DNA to Cyanobacteria. *Methods in Enzymology* 167, 747-54) The liquid cultures were then assayed for tocopherols by harvesting 1ml of culture by centrifugation,

extracting with ethanol/pyrogallol, and HPLC separation. The slr1736 *Synechocystis 6803* KO strain, did not contain any detectable tocopherols, while the slr1736 *Synechocystis 6803* KO strain transformed with pmon21690 contained detectable alpha tocopherol. A *Synechocystis 6803* strain transformed with psl1211(vector control) produced alpha tocopherol as well.

5

**Example 5:** Transgenic Plant Analysis

Arabidopsis plants transformed with constructs for the sense or antisense expression of the ATPT proteins were analyzed by High Pressure Liquid Chromatography (HPLC) for altered levels of total tocopherols, as well as altered levels of specific tocopherols (alpha, beta, gamma, and delta tocopherol).

Extracts of leaves and seeds were prepared for HPLC as follows. For seed extracts, 10 mg of seed was added to 1 g of microbeads (Biospec) in a sterile microfuge tube to which 500 ul 1% pyrogallol (Sigma Chem)/ethanol was added. The mixture was shaken for 3 minutes in a mini Beadbeater (Biospec) on “fast” speed. The extract was filtered through a 0.2 um filter into an autosampler tube. The filtered extracts were then used in HPLC analysis described below.

Leaf extracts were prepared by mixing 30-50 mg of leaf tissue with 1 g microbeads and freezing in liquid nitrogen until extraction. For extraction, 500 ul 1% pyrogallol in ethanol was added to the leaf/bead mixture and shaken for 1 minute on a Beadbeater (Biospec) on “fast” speed. The resulting mixture was centrifuged for 4 minutes at 14,000 rpm and filtered as described above prior to HPLC analysis.

HPLC was performed on a Zorbax silica HPLC column (4.6 mm X 250 mm) with a fluorescent detection, an excitation at 290 nm, an emission at 336 nm, and bandpass and slits. Solvent A was hexane and solvent B was methyl-t-butyl ether. The injection volume was 20 ul, the flow rate was 1.5 ml/min, the run time was 12 min (40°C) using the gradient (Table 5):

**Table 5:**

	<u>Time</u>	<u>Solvent A</u>	<u>Solvent B</u>
30	0 min.	90%	10%
	10 min.	90%	10%
	11 min.	25%	75%
	12 min.	90%	10%

Tocopherol standards in 1% pyrogallol/ ethanol were also run for comparison (alpha tocopherol, gamma tocopherol, beta tocopherol, delta tocopherol, and tocopherol (tocol) (all from Matreya).

5 Standard curves for alpha, beta, delta, and gamma tocopherol were calculated using Chemstation software. The absolute amount of component x is: Absolute amount of x= Response<sub>x</sub> x RF<sub>x</sub> x dilution factor where Response<sub>x</sub> is the area of peak x, RF<sub>x</sub> is the response factor for component x (Amount<sub>x</sub>/Response<sub>x</sub>) and the dilution factor is 500 ul. The ng/mg tissue is found by: total ng component/mg plant tissue.

10 Results of the HPLC analysis of seed extracts of transgenic *Arabidopsis* lines containing pMON10822 for the expression of ATAT2 from the napin promoter are provided in Figure 24.

HPLC analysis results of *Arabidopsis* seed tissue expressing the ATAT2 sequence from the napin promoter (pMON10822) demonstrates an increased level of tocopherols in the seed. Total tocopherol levels are increased as much as 50 to 60% over the total tocopherol levels of non-transformed (wild-type) *Arabidopsis* plants (Figure 24).

Furthermore, increases of particular tocopherols are also increased in transgenic *Arabidopsis* plants expressing the ATAT2 nucleic acid sequence from the napin promoter. Levels of delta tocopherol in these lines are increased greater than 3 fold over the delta tocopherol levels obtained from the seeds of wild type *Arabidopsis* lines. Levels of gamma tocopherol in transgenic *Arabidopsis* lines expressing the ATAT2 nucleic acid sequence are increased as much as about 60% over the levels obtained in the seeds of non-transgenic control lines. Furthermore, levels of alpha tocopherol are increased as much as 3 fold over those obtained from non-transgenic control lines.

25 Results of the HPLC analysis of seed extracts of transgenic *Arabidopsis* lines containing pMON10803 for the expression of ATAT2 from the enhanced 35S promoter are provided in Figure 25.

30 All publications and patent applications mentioned in this specification are indicative of the level of skill of those skilled in the art to which this invention pertains. All publications and

patent applications are herein incorporated by reference to the same extent as if each individual publication or patent application was specifically and individually indicated to be incorporated by reference.

Although the foregoing invention has been described in some detail by way of illustration  
5 and example for purposes of clarity of understanding, it will be obvious that certain changes and modifications may be practiced within the scope of the appended claim.

DRAFT - DO NOT CITE OR RELY UPON

## Claims

What is Claimed is:

1. An isolated nucleic acid sequence encoding a prenyltransferase.
- 5 2. An isolated nucleic acid sequence according to Claim 1, wherein said prenyltransferase is selected from the group consisting of straight chain prenyltransferase and aromatic prenyltransferase.
3. An isolated DNA sequence according to Claim 1, wherein said nucleic acid sequence is isolated from a eukaryotic cell source.
- 10 4. An isolated DNA sequence according to Claim 3, wherein said eukaryotic cell source is selected from the group consisting of mammalian, nematode, fungal, and plant cells.
5. The DNA encoding sequence of Claim 4 wherein said prenyltransferase protein is from *Arabidopsis*.
6. The DNA encoding sequence of Claim 5 wherein said prenyltransferase protein is encoded by a sequence selected from the group consisting of the sequences of Figure 1.
7. The DNA encoding sequence of Claim 4 wherein said prenyltransferase protein is from corn.
8. The DNA encoding sequence of Claim 7 wherein said prenyltransferase protein is encoded by a sequence which includes the EST of the sequences of Figure 3.
9. The DNA encoding sequence of Claim 4 wherein said prenyltransferase protein is from soybean.
10. The DNA encoding sequence of Claim 9 wherein said prenyltransferase protein is encoded by a sequence which includes the ESTs of the group consisting of the sequences of Figure 2 and Figure 9.
11. An isolated DNA sequence according to Claim 1, wherein said nucleic acid sequence is isolated from a prokaryotic cell source.
- 25 12. An isolated DNA sequence according to Claim 11, wherein said prokaryotic source is *Synechocystis*.
13. A nucleic acid construct comprising as operably linked components, a transcriptional initiation region functional in a host cell, a nucleic acid sequence encoding prenyltransferase, and a transcriptional termination region.

14. A nucleic acid construct according to Claim 13, wherein said nucleic acid sequence encoding prenyltransferase is obtained from an organism selected from the group consisting of a eukaryotic organism and a prokaryotic organism.

5 15. A nucleic acid construct according to Claim 14, wherein said nucleic acid sequence encoding prenyltransferase is obtained from a plant source.

16. A nucleic acid construct according to Claim 15, wherein said nucleic acid sequence encoding prenyltransferase is obtained from a source selected from the group consisting of *Arabidopsis*, soybean and corn.

10 17. A nucleic acid construct according to Claim 13, wherein said nucleic acid sequence encoding prenyltransferase is obtained from *Synechocystis*.

18. A plant cell comprising the construct of Claim 13.

19. A method for the alteration of the tocopherol content in a host cell, comprising; transforming said host cell with a construct comprising as operably linked components, a transcriptional initiation region functional in a host cell, a nucleic acid sequence encoding prenyltransferase, and a transcriptional termination region.

20. The method according to Claim 19, wherein said host cell is selected from the group consisting of a prokaryotic cell and a eukaryotic cell.

21. The method according to Claim 20, wherein said prokaryotic cell is *Synechocystis*.

22. The method according to Claim 20, wherein said eukaryotic cell is a plant cell.

23. The method according to Claim 22, wherein said plant cell is obtained from a plant selected from the group consisting of *Arabidopsis*, soybean, and corn.

24. A method for producing a tocopherol compound of interest in a host cell, said method comprising obtaining a transformed host cell, said host cell having and expressing in its genome: a construct having a DNA sequence encoding a prenyltransferase operably linked to a transcriptional initiation region functional in a host cell,

25 wherein said prenyltransferase is involved in the synthesis of tocopherols.

26. The method according to Claim 24, wherein said host cell is selected from the group consisting of a prokaryotic cell and a eukaryotic cell.

27. The method according to Claim 25, wherein said prokaryotic cell is *Synechocystis*.

30 28. The method according to Claim 24, wherein said eukaryotic cell is a plant cell.

28. The method according to Claim 27, wherein said plant cell is obtained from a plant selected from the group consisting of *Arabidopsis*, soybean, and corn.
29. A method for increasing the biosynthetic flux in cell from a host cell toward tocopherol production, said method comprising transforming said host cell with a construct comprising as operably linked components, a transcriptional initiation region functional in a host cell, a DNA encoding a prenyltransferase involved in the synthesis of tocopherols, and a transcriptional termination region.
30. The method according to Claim 29, wherein said host cell is selected from the group consisting of a prokaryotic cell and a eukaryotic cell.
31. The method according to Claim 30, wherein said prokaryotic cell is *Synechocystis*.
32. The method according to Claim 30, wherein said eukaryotic cell is a plant cell.
33. The method according to Claim 32, wherein said plant cell is obtained from a plant selected from the group consisting of *Arabidopsis*, soybean, and corn.

17133/01/US

## NUCLEIC ACID SEQUENCES TO PROTEINS INVOLVED IN TOCOPHEROL SYNTHESIS

5

### Abstract

Nucleic acid sequences and methods are provided for producing plants and seeds having altered tocopherol content and compositions. The methods find particular use in increasing the tocopherol levels in plants, and in providing desirable tocopherol compositions in a host plant 10 cell.

DRAFTING IN PROGRESS

ATPT2 : \* 20 \* 40 \* 60 \* 80 \*  
ATPT3 : \* 20 \* 40 \* 60 \* 80 \*  
ATPT4 : \* 20 \* 40 \* 60 \* 80 \*  
ATPT8 : \* 20 \* 40 \* 60 \* 80 \*  
ATPT12 : \* 20 \* 40 \* 60 \* 80 \*

ATPT2 : \* 20 \* 40 \* 60 \* 80 \*  
ATPT3 : \* 20 \* 40 \* 60 \* 80 \*  
ATPT4 : \* 20 \* 40 \* 60 \* 80 \*  
ATPT8 : \* 20 \* 40 \* 60 \* 80 \*  
ATPT12 : \* 20 \* 40 \* 60 \* 80 \*  
  
ATPT2 : SSSLVY\* 100 \* 120 \* 140 \* 160 \* 180 \*  
ATPT3 : SSSLVY\* 100 \* 120 \* 140 \* 160 \* 180 \*  
ATPT4 : SSSLVY\* 100 \* 120 \* 140 \* 160 \* 180 \*  
ATPT8 : SSSLVY\* 100 \* 120 \* 140 \* 160 \* 180 \*  
ATPT12 : SSSLVY\* 100 \* 120 \* 140 \* 160 \* 180 \*  
  
ATPT2 : SSSLVY\* 200 \* 220 \* 240 \* 260 \* 280 \*  
ATPT3 : SSSLVY\* 200 \* 220 \* 240 \* 260 \* 280 \*  
ATPT4 : SSSLVY\* 200 \* 220 \* 240 \* 260 \* 280 \*  
ATPT8 : SSSLVY\* 200 \* 220 \* 240 \* 260 \* 280 \*  
ATPT12 : SSSLVY\* 200 \* 220 \* 240 \* 260 \* 280 \*  
  
ATPT2 : VAAALMNIIVGQ\* 300 \* 320 \* 340 \* 360 \*  
ATPT3 : VAAALMNIIVGQ\* 300 \* 320 \* 340 \* 360 \*  
ATPT4 : VAAALMNIIVGQ\* 300 \* 320 \* 340 \* 360 \*  
ATPT8 : VAAALMNIIVGQ\* 300 \* 320 \* 340 \* 360 \*  
ATPT12 : VAAALMNIIVGQ\* 300 \* 320 \* 340 \* 360 \*  
  
ATPT2 : PLRWRKRFALVAAMCIAVRAILYQTAFYTHIOTHYFGRPLFLTRPLIFATAFMSFFS\* 400 \* 420 \* 440 \*  
ATPT3 : B-LMKRFTFWPOAFLIGHT\* 400 \* 420 \* 440 \*  
ATPT4 : LKQLHPINTWVGAVN\* 400 \* 420 \* 440 \*  
ATPT8 : LAFEGYGRNLGAFOQI\* 400 \* 420 \* 440 \*  
ATPT12 : PDKLKGONGWGNFA\* 400 \* 420 \* 440 \*  
  
ATPT2 : FCTRS\* 460 \* 480 \*  
ATPT3 : FCTRS\* 460 \* 480 \*  
ATPT4 : FCTRS\* 460 \* 480 \*  
ATPT8 : FCTRS\* 460 \* 480 \*  
ATPT12 : FCTRS\* 460 \* 480 \*

Figure 1

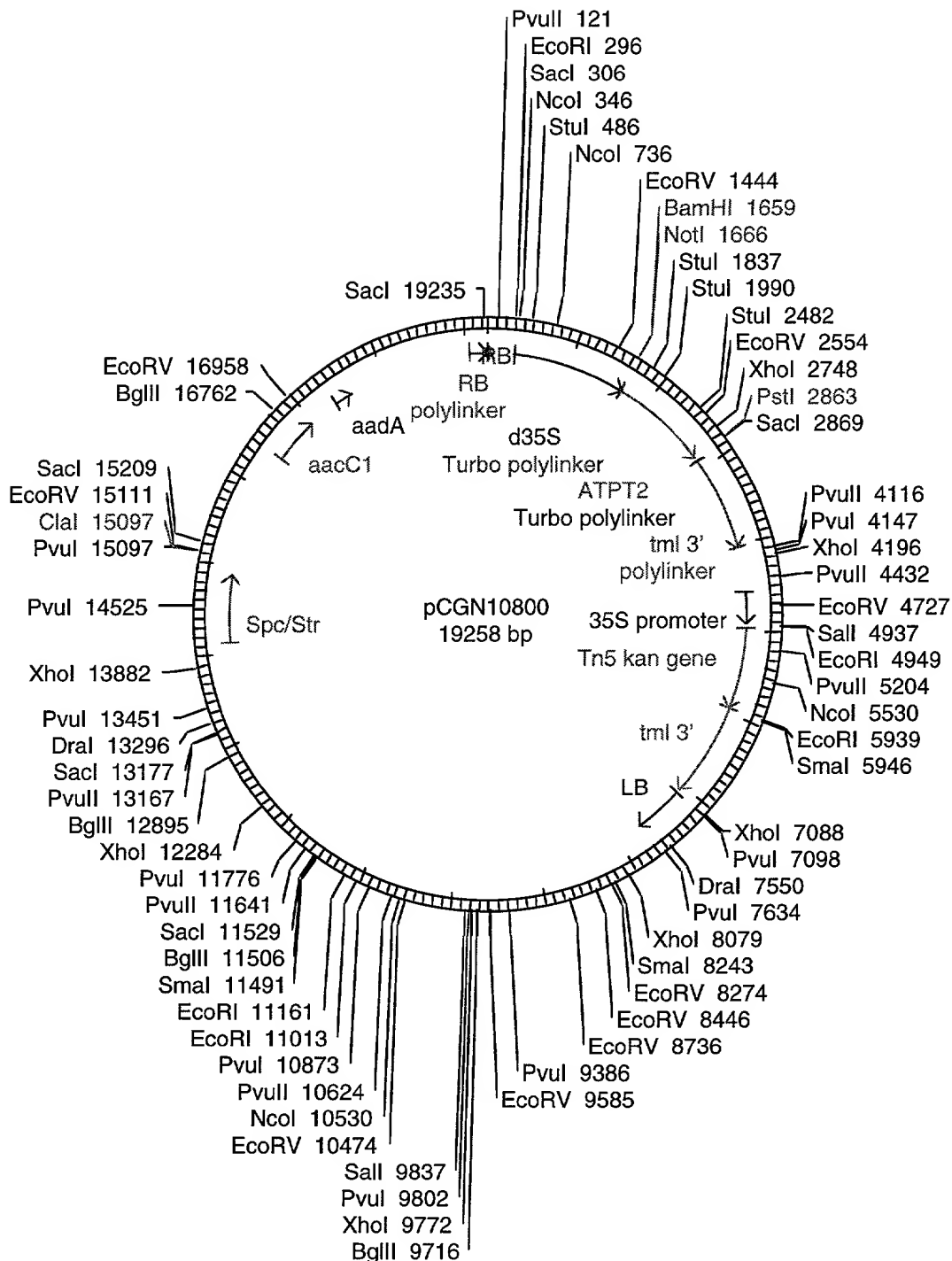


Figure 2

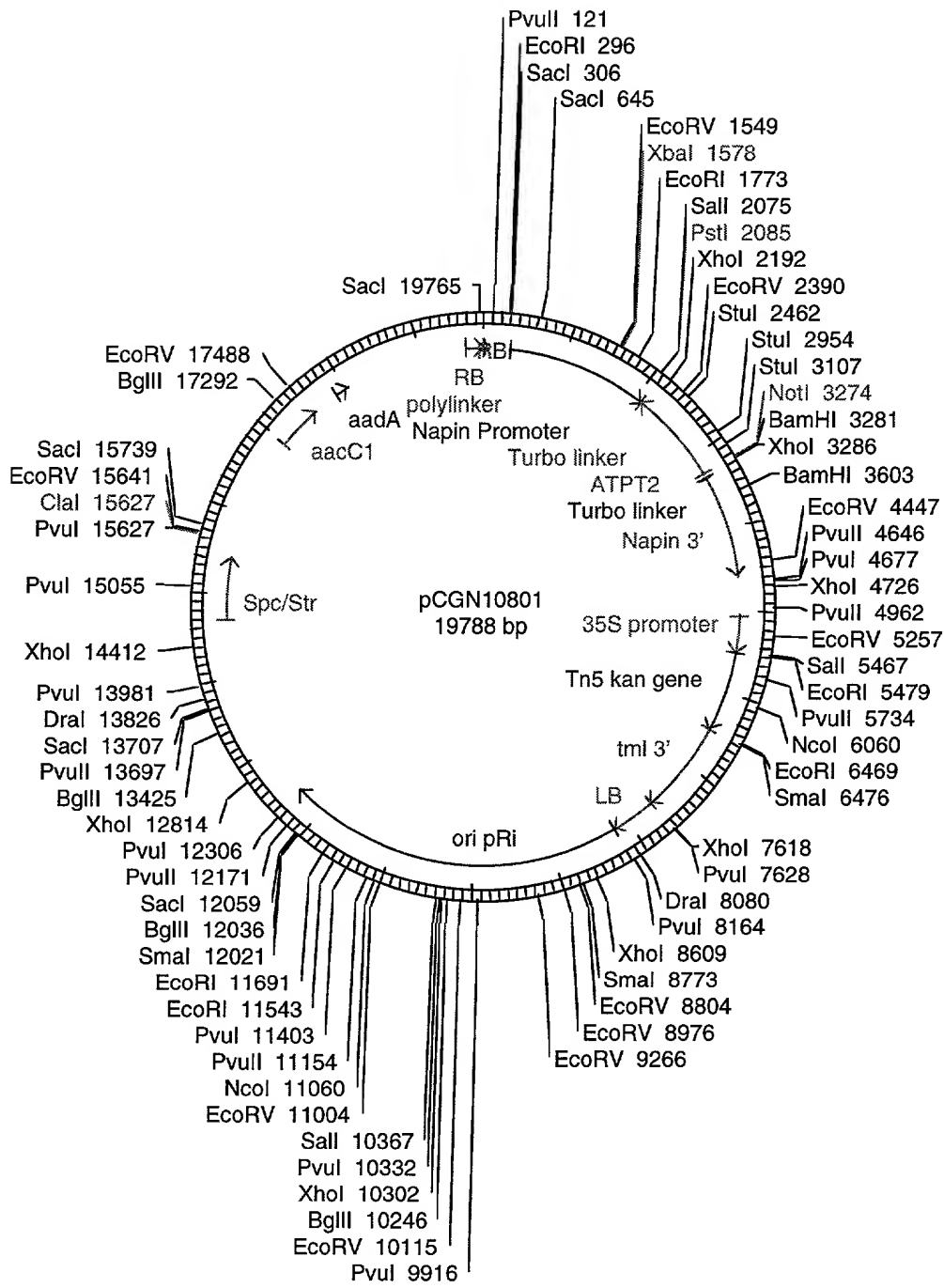


Figure 3

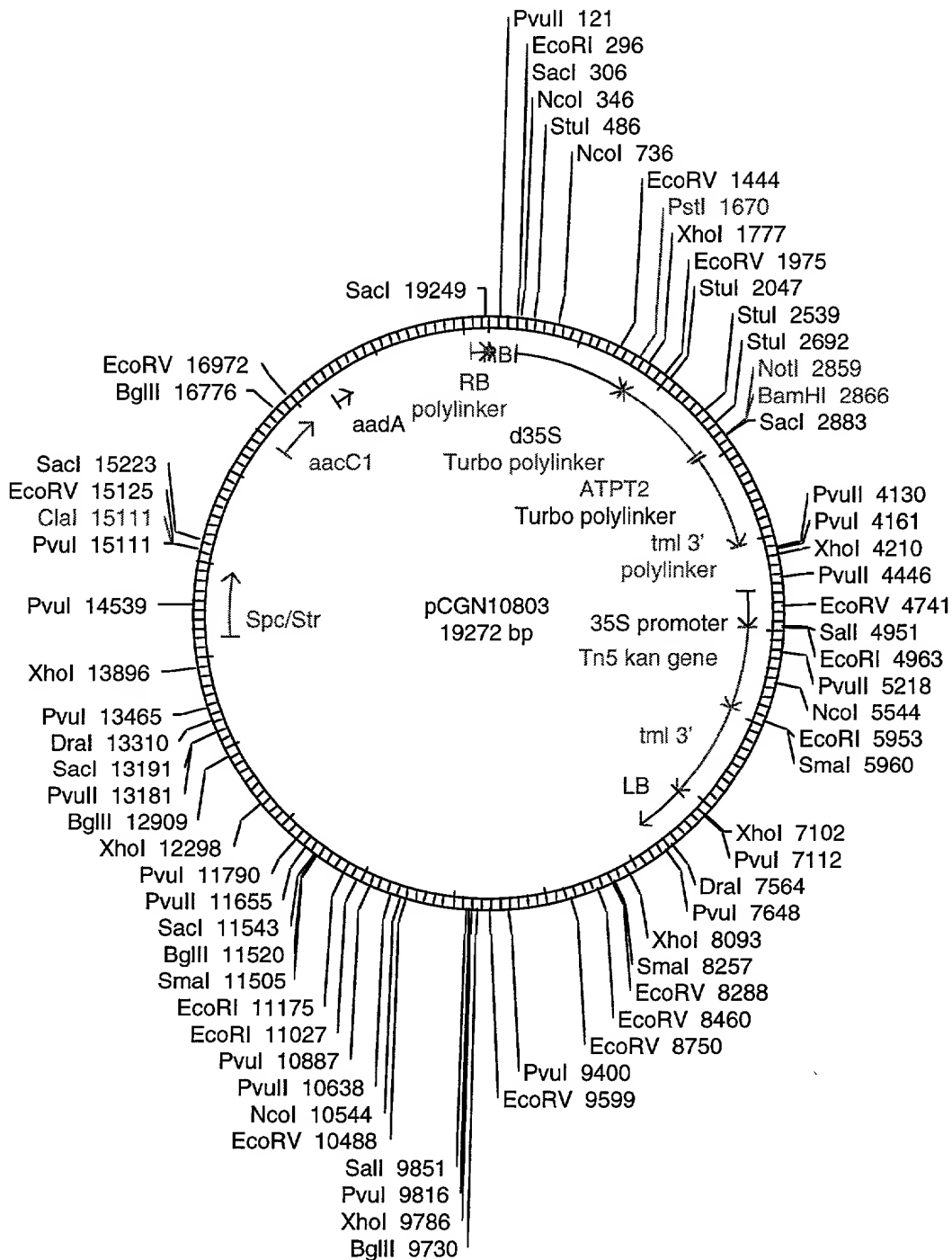


Figure 4

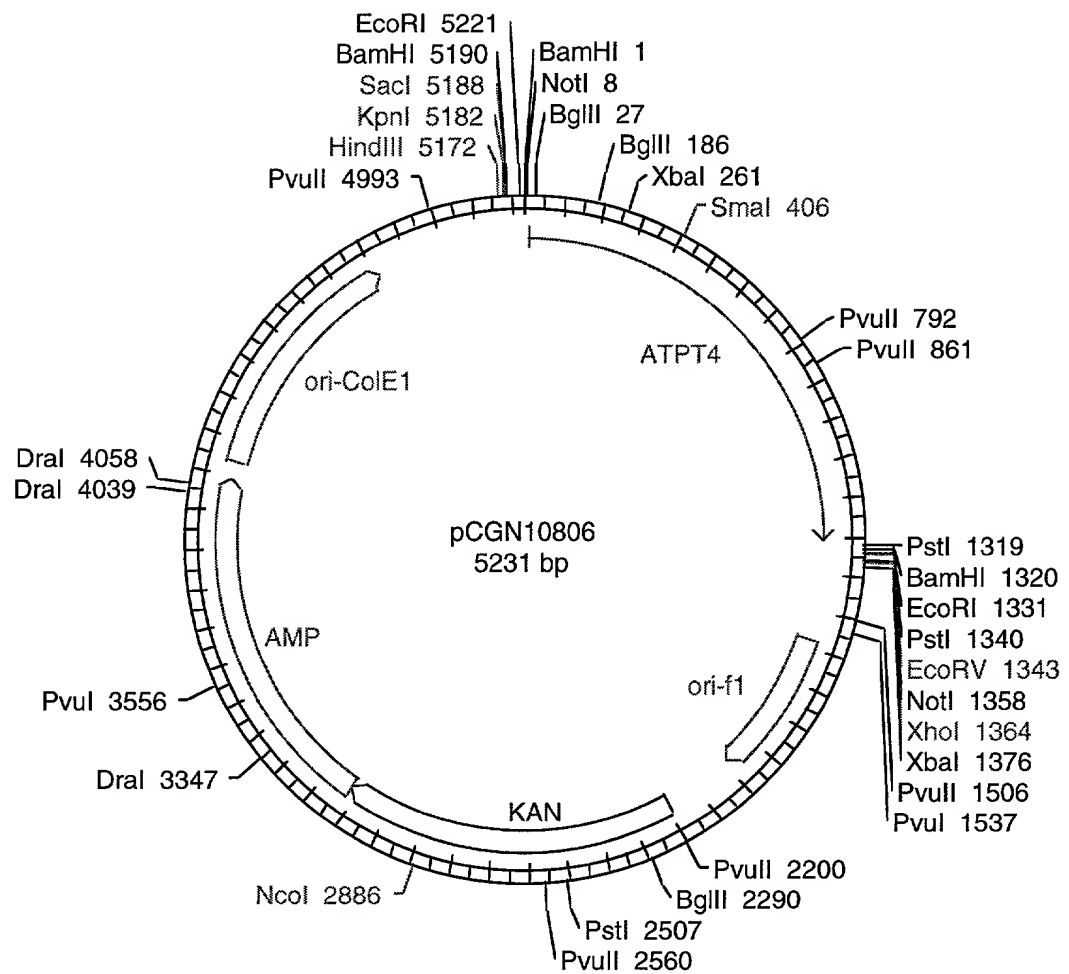


Figure 5

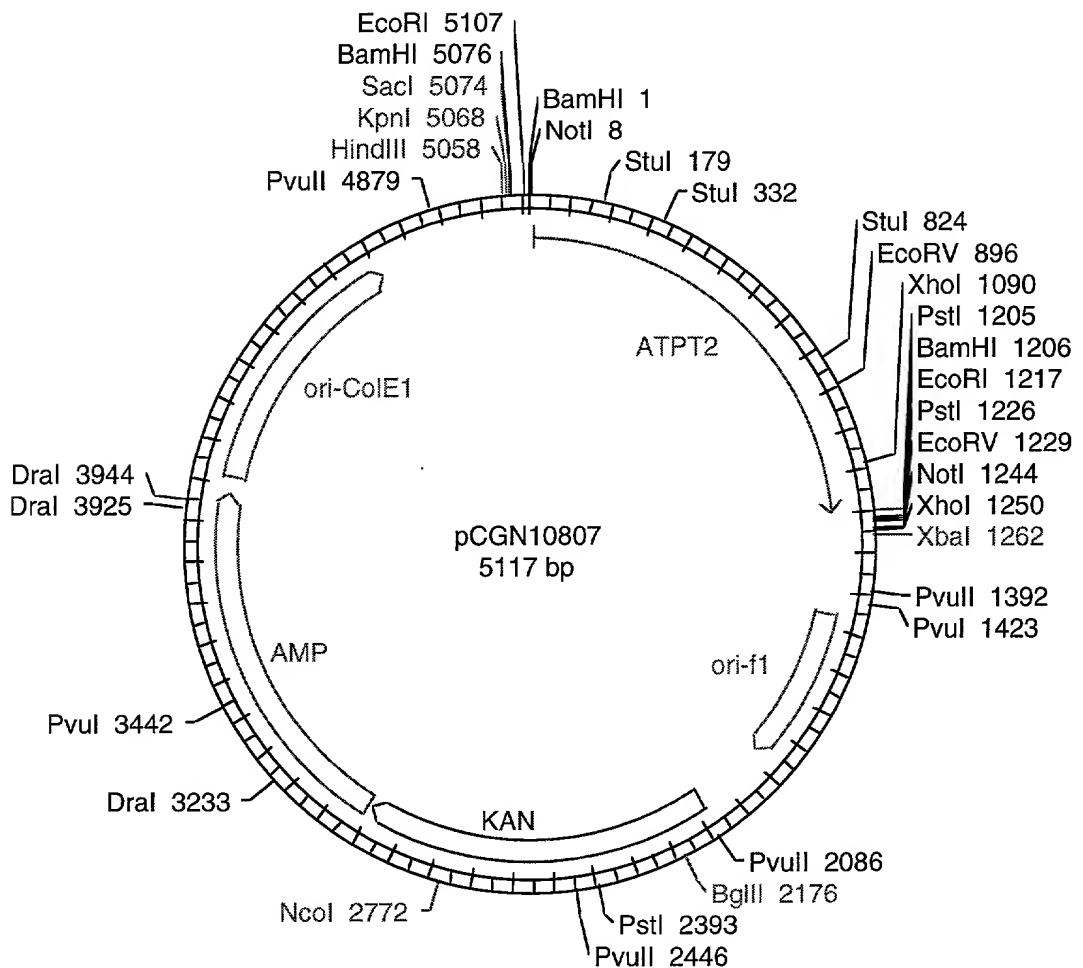


Figure 6

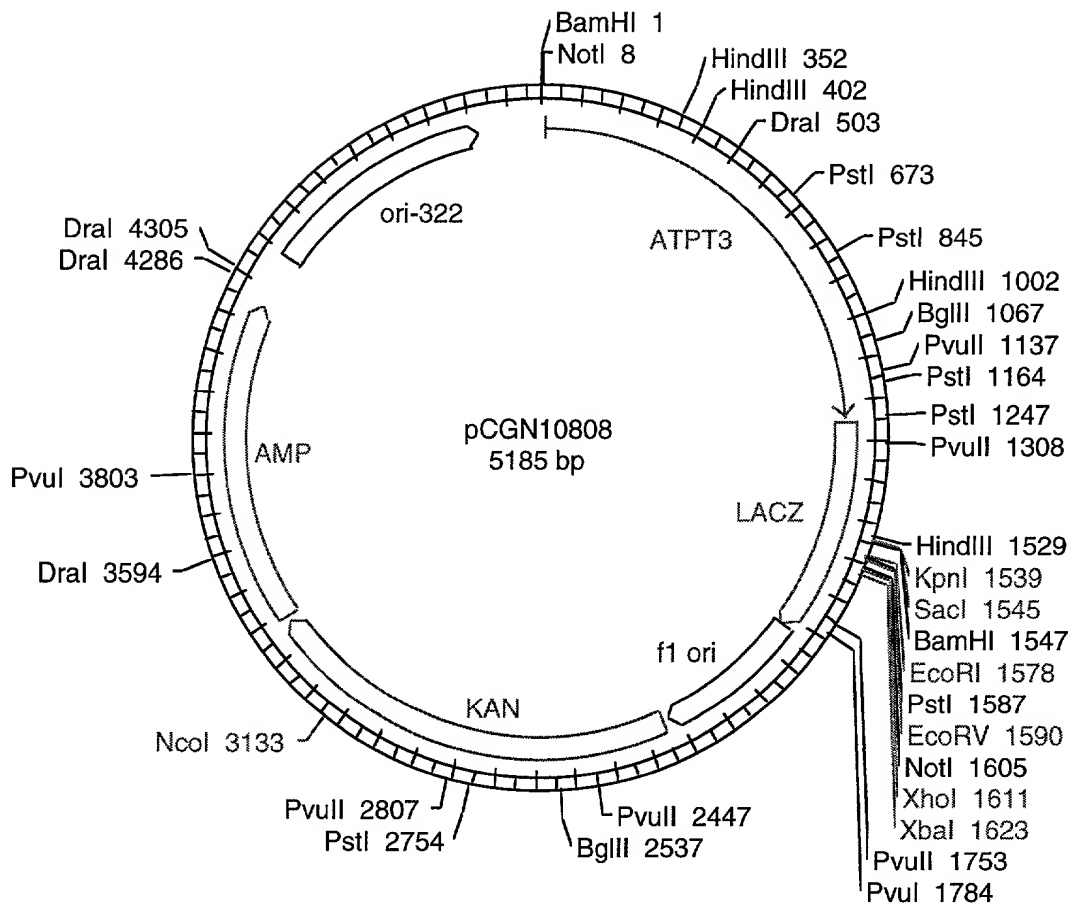


Figure 7

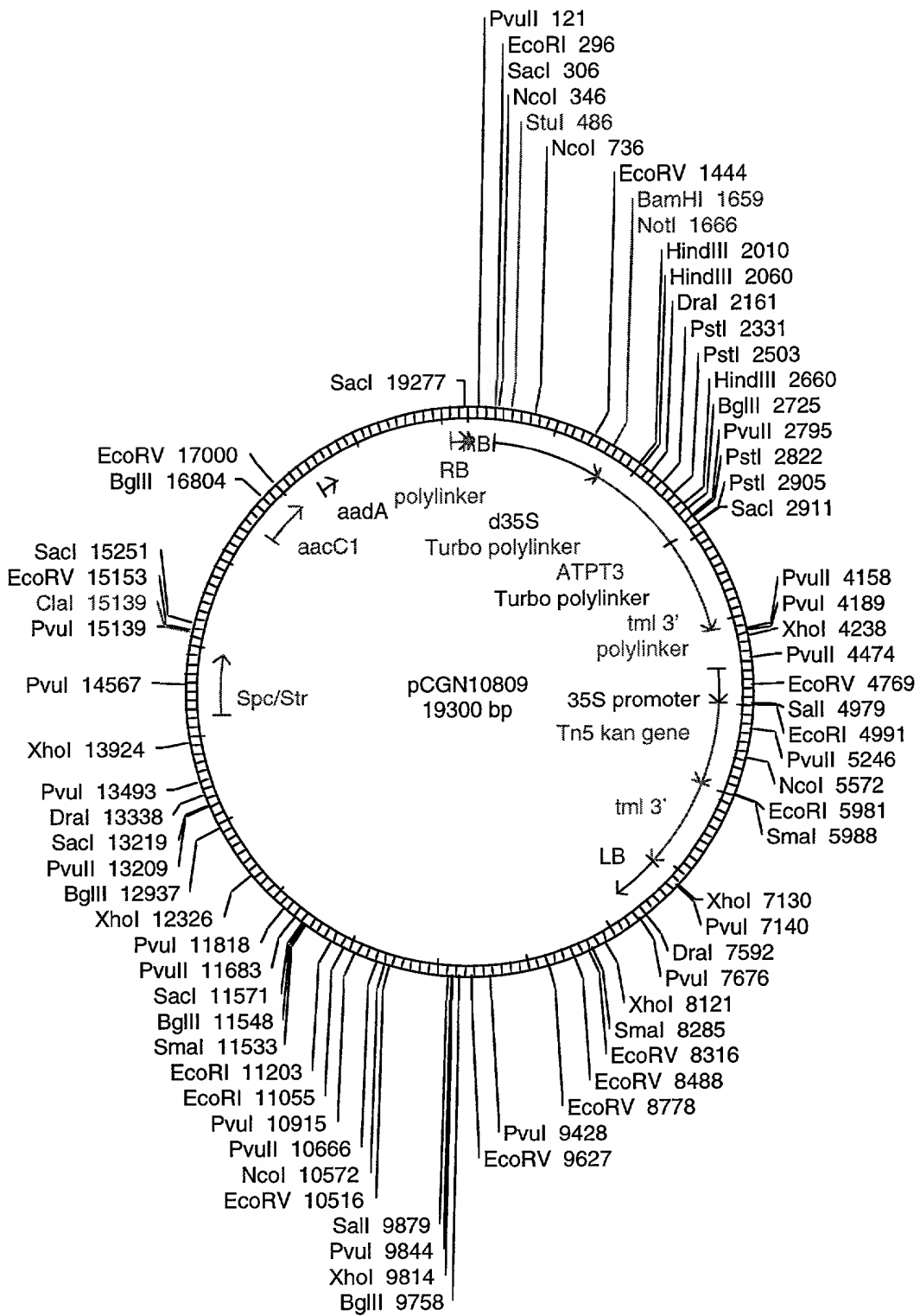


Figure 8

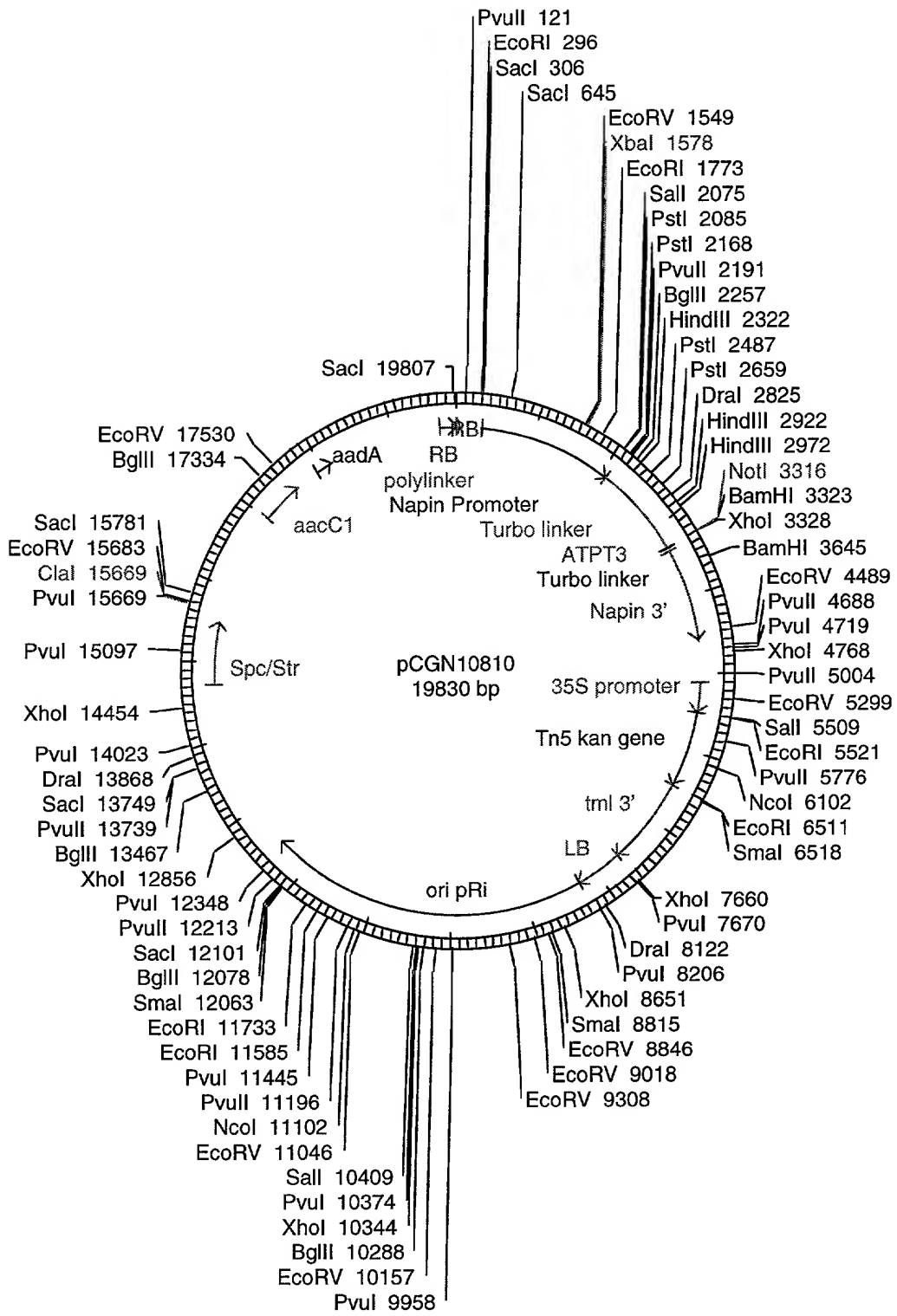


Figure 9

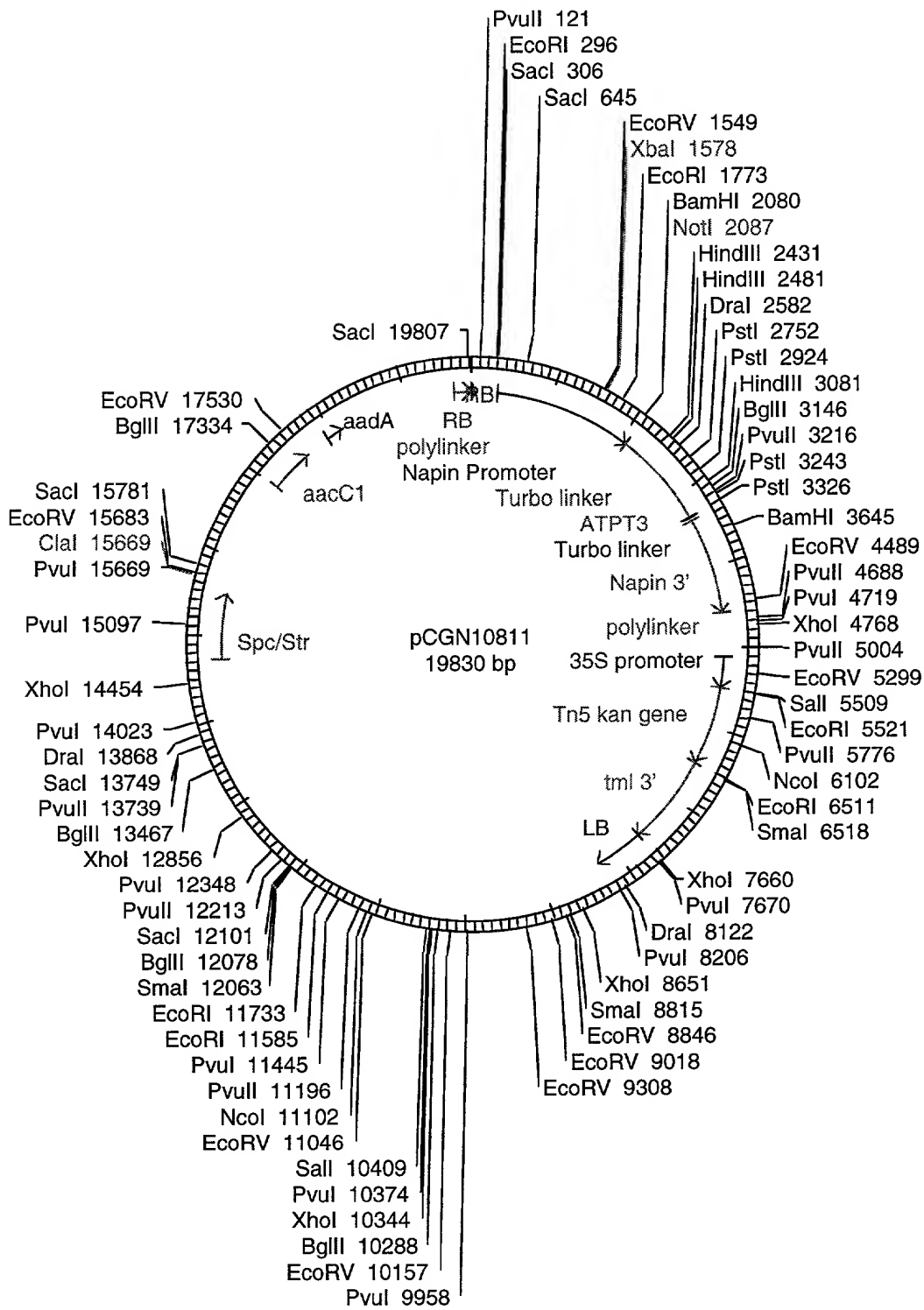


Figure 10

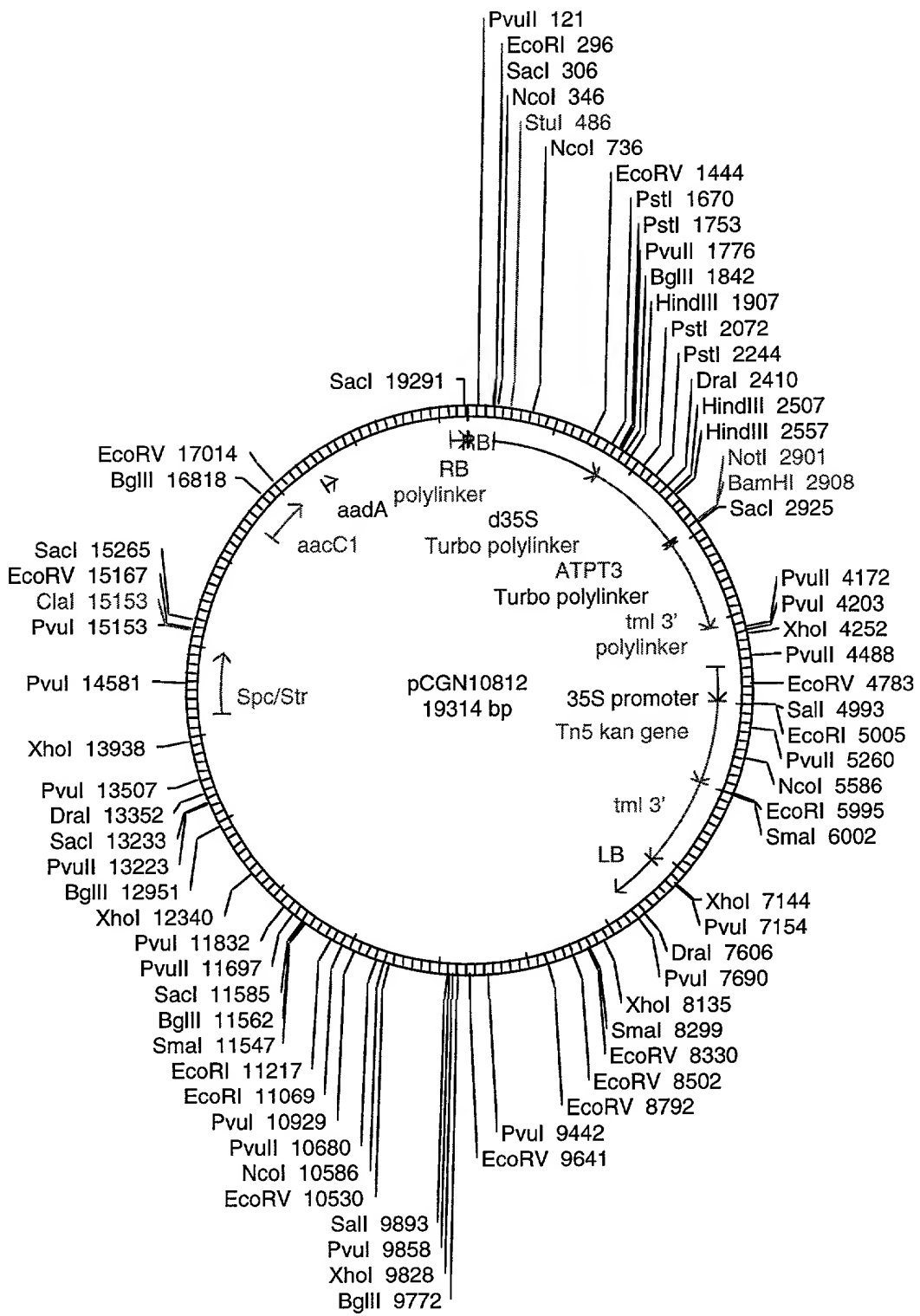


Figure 11

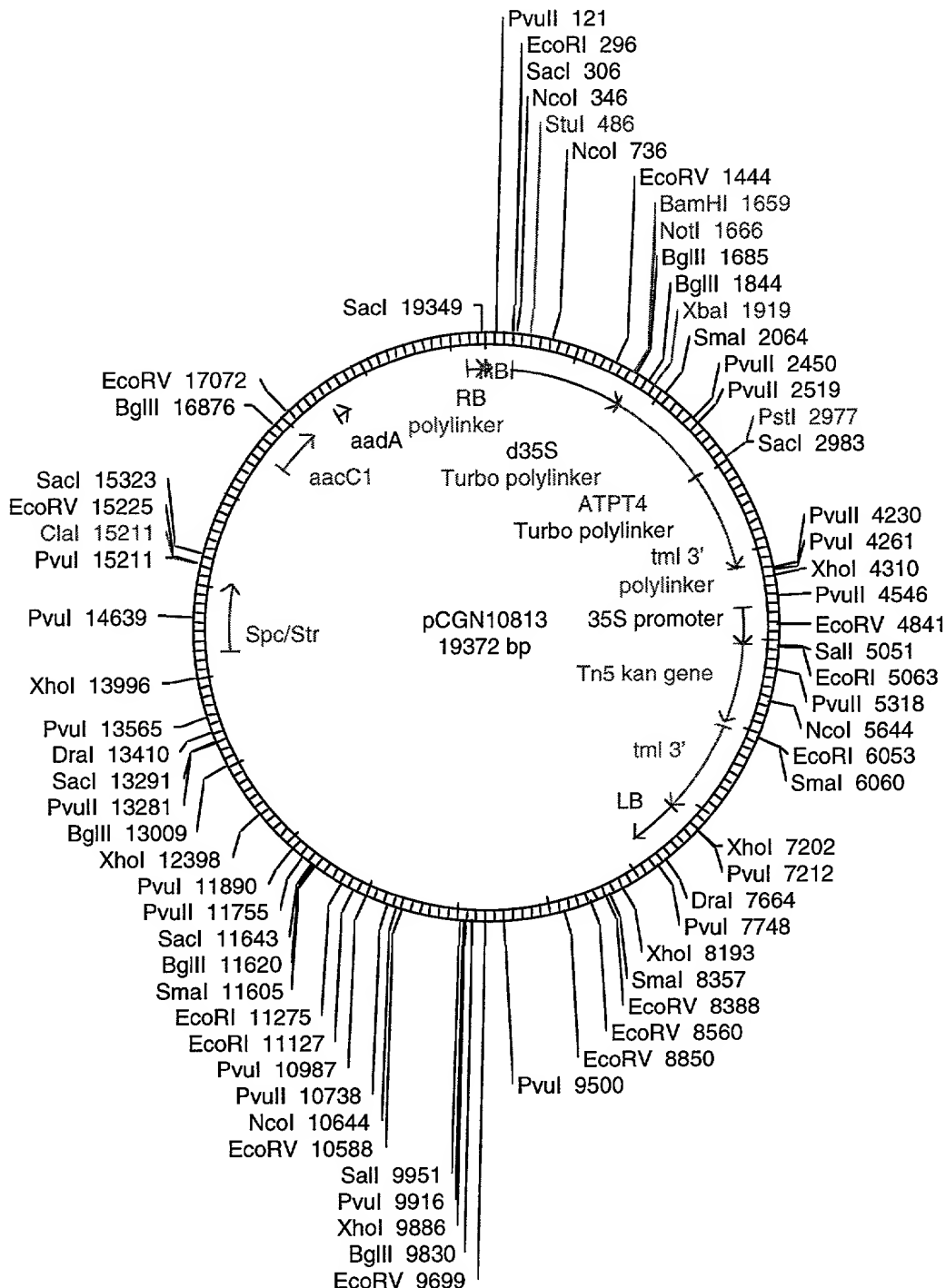


Figure 12

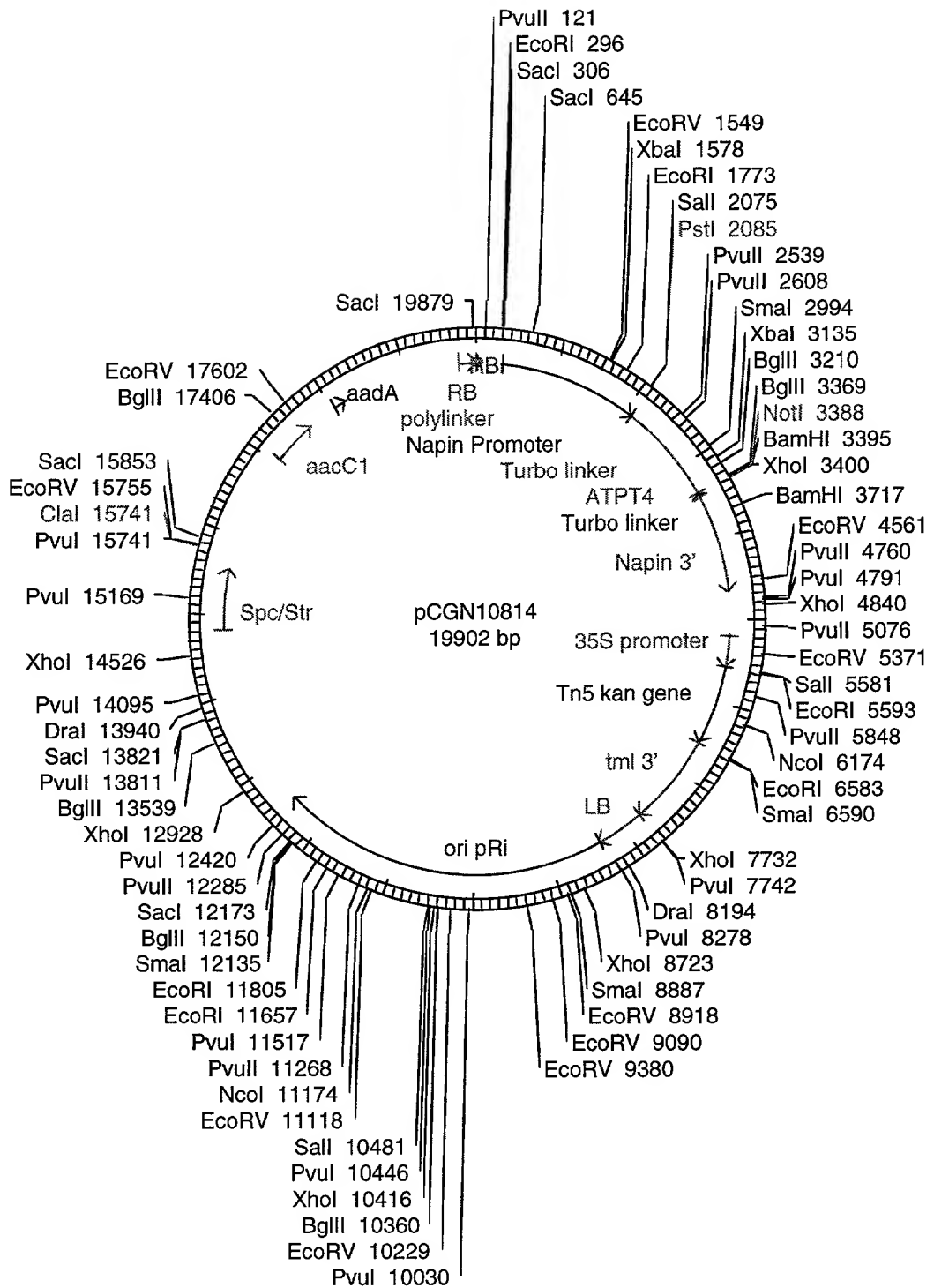


Figure 13

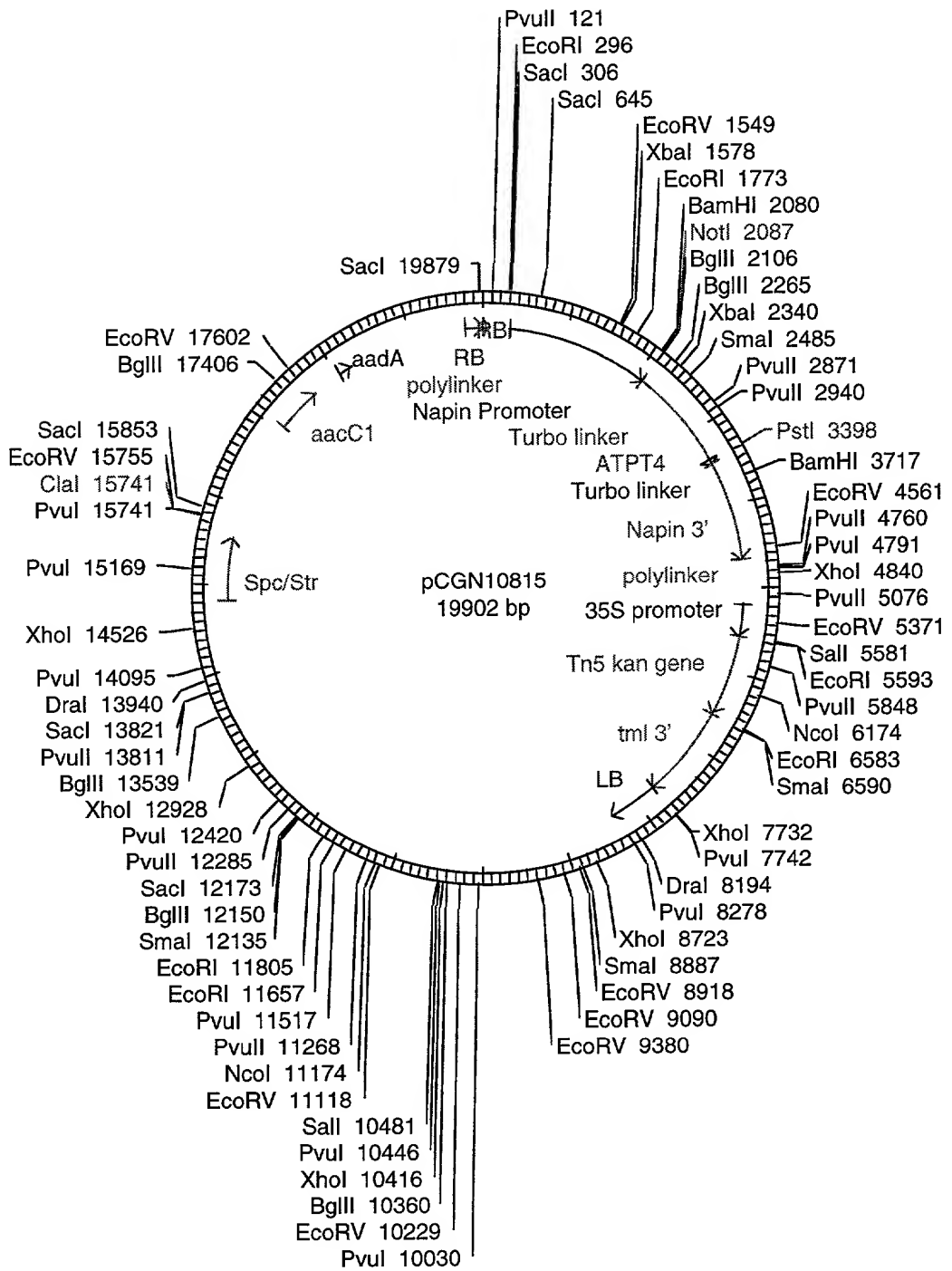


Figure 14

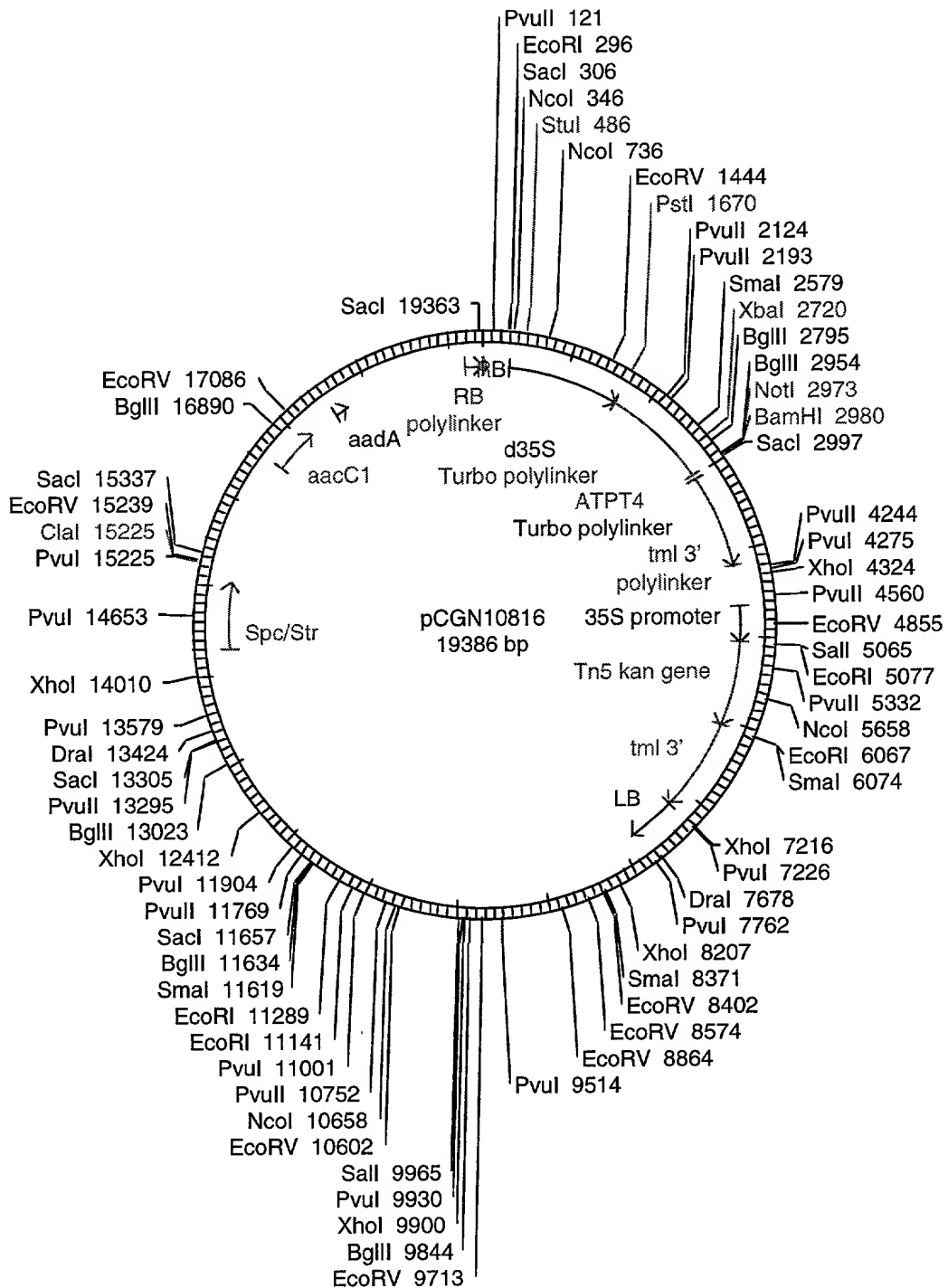


Figure 15

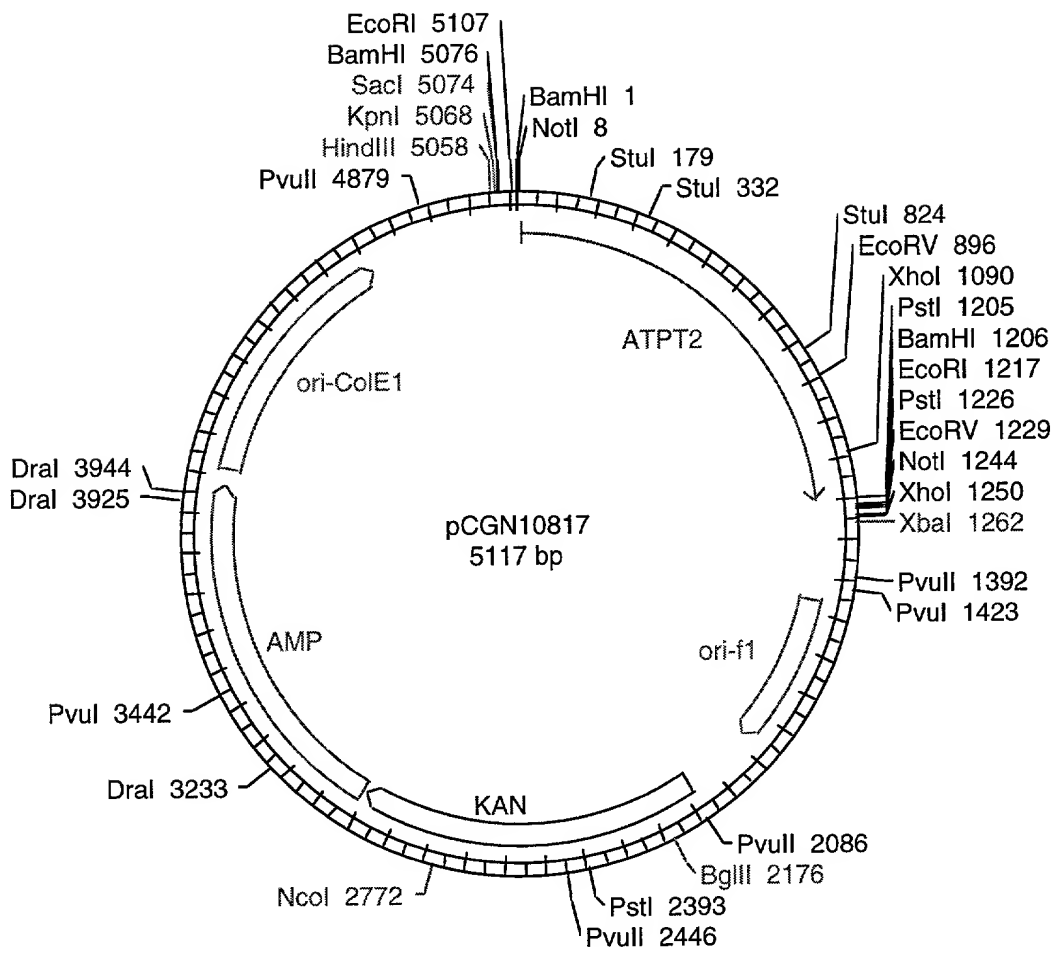


Figure 16

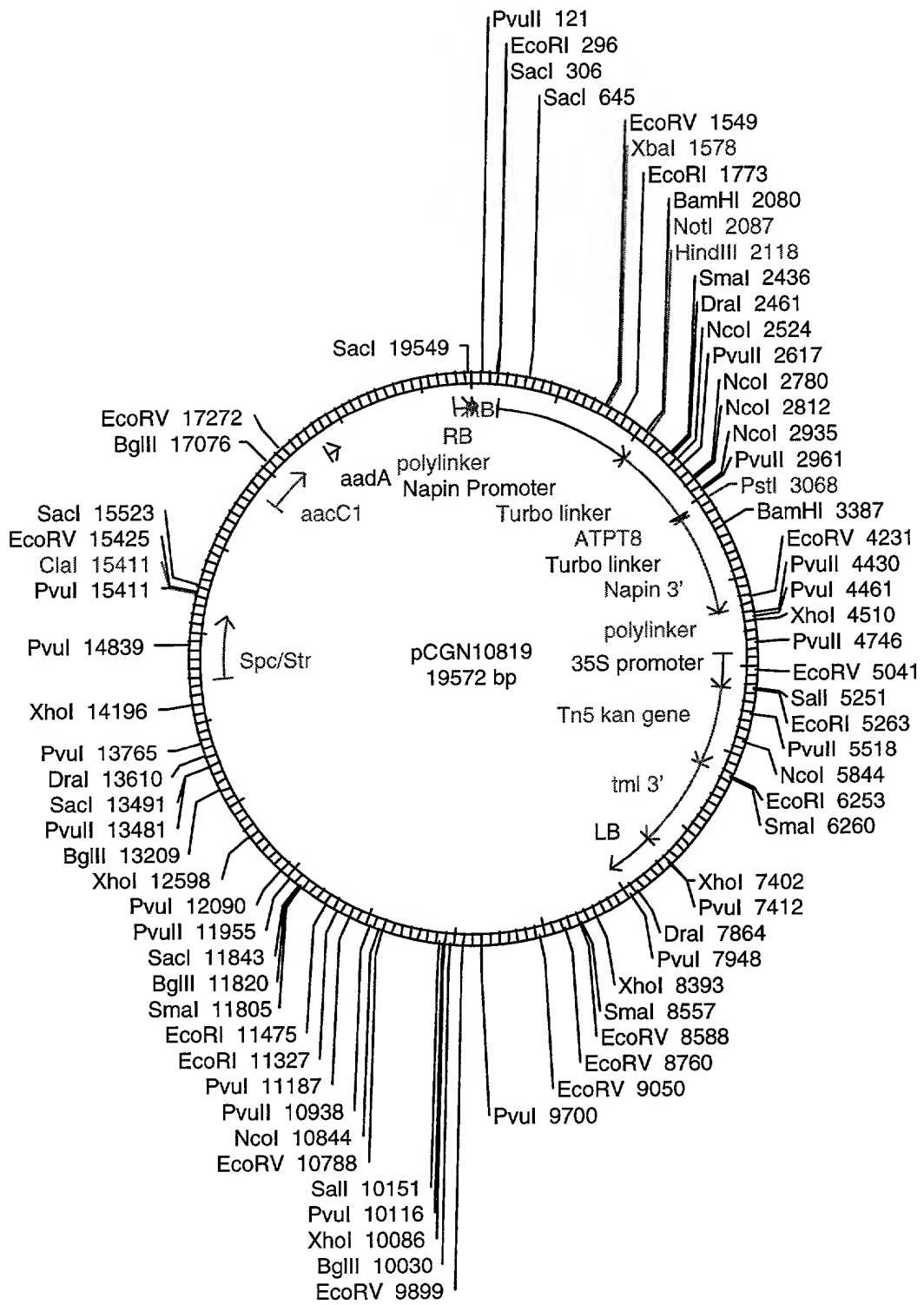


Figure 17

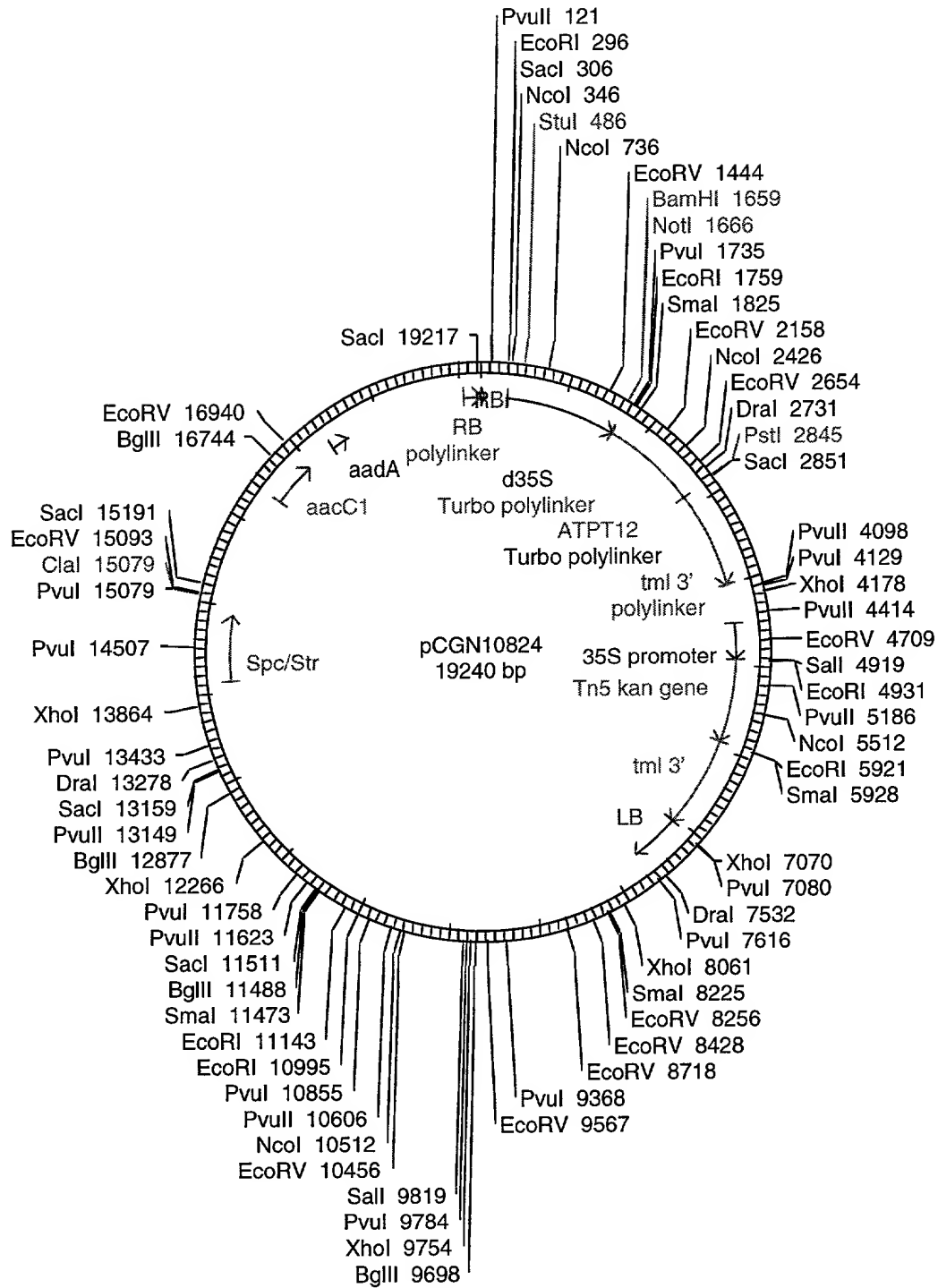


Figure 18

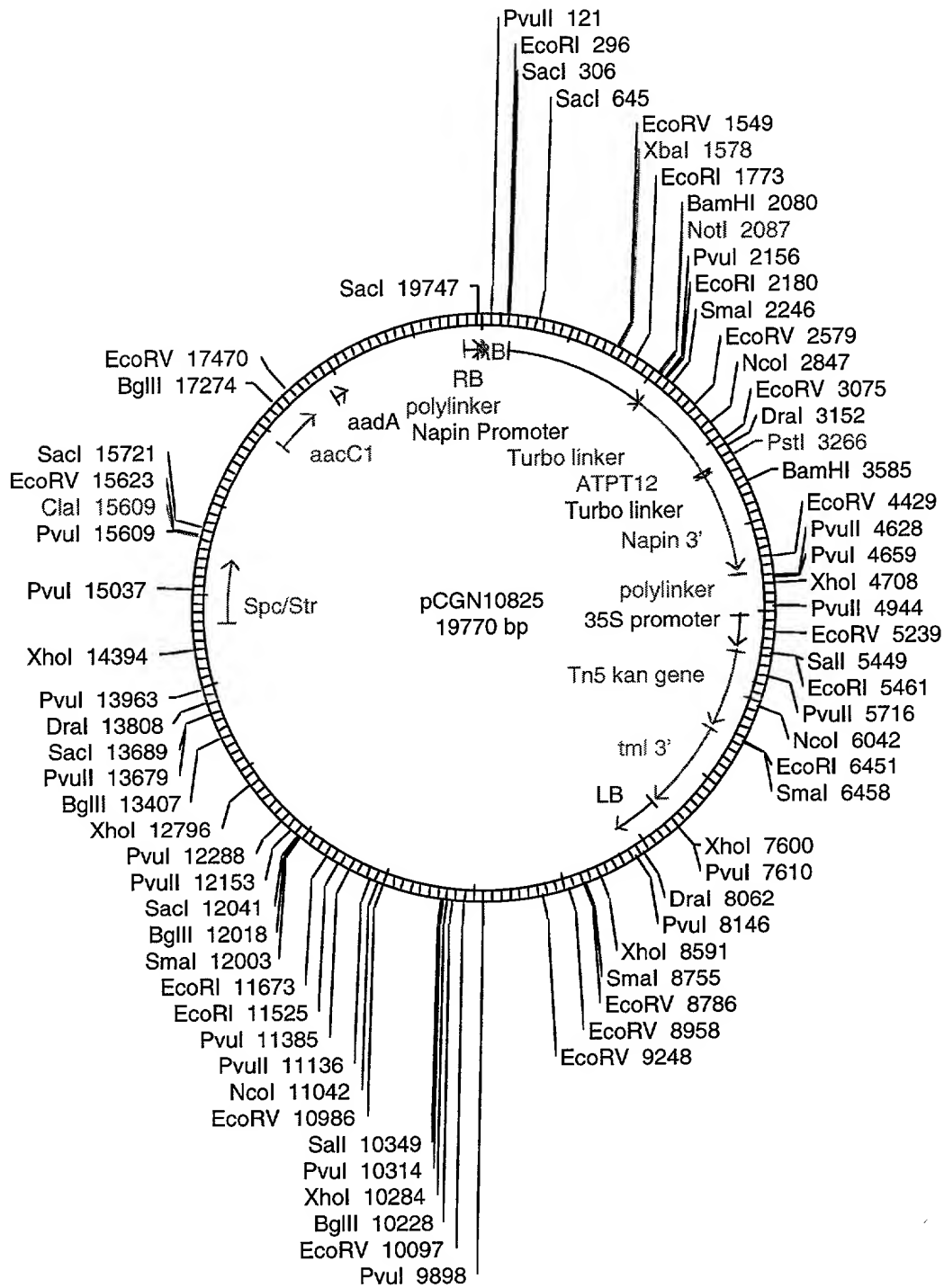


Figure 19

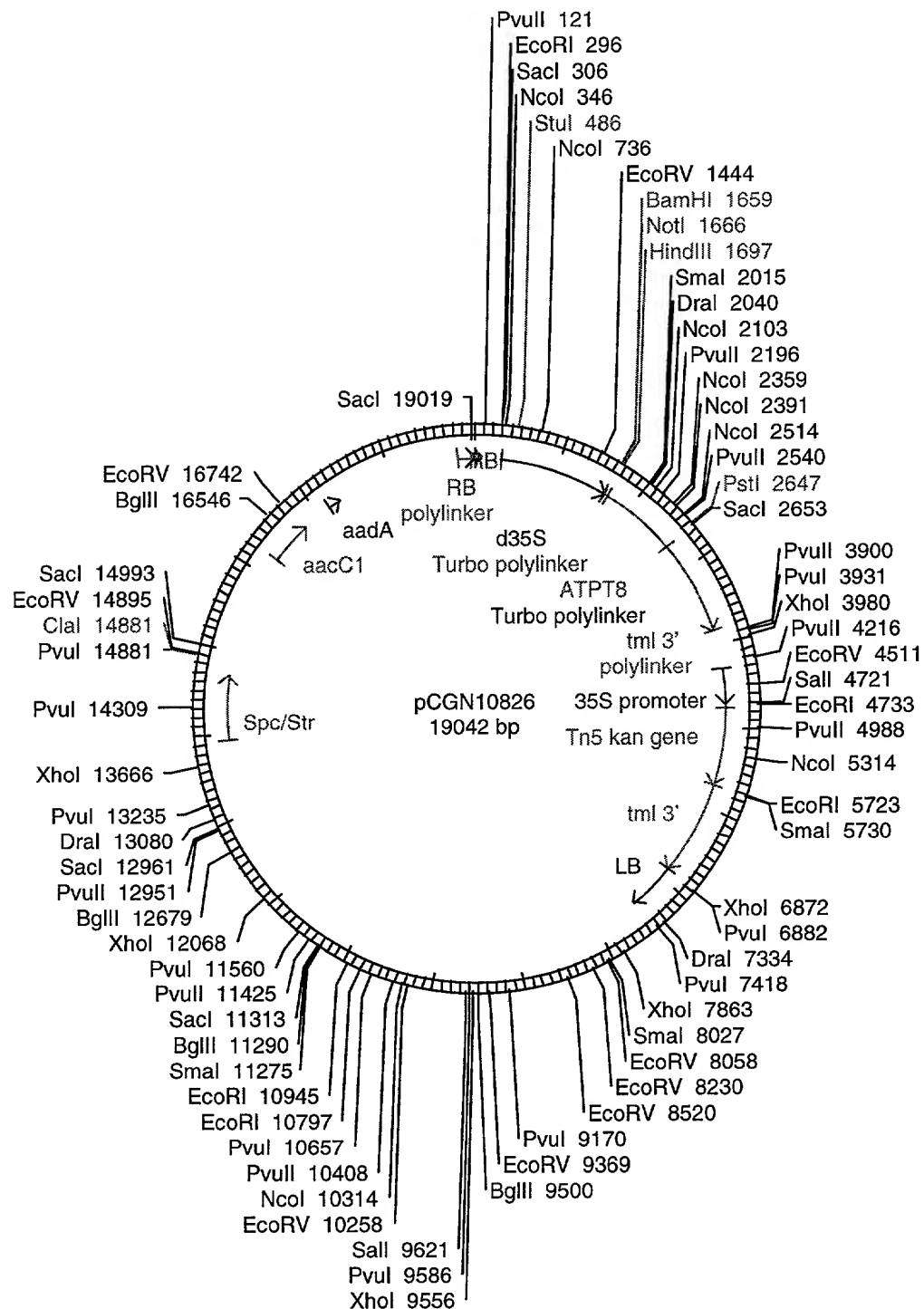


Figure 20

CHARTERED

SLR1736 :	MATIOAFWRR-----FSRPHTI	GTTLS-----WAVYL	TILGD--G-NSVNSPAS	* 60	* 80	*
SLR0926 :	MVAQTPSS---PPLWLTLLYLRWHEK	-----AGRLLEM	BALWA-----CLAAQELPP	-----	DLYFGAWL	IVG-----*
SLL1899 :	MVTSTKIHQRHDSMGAVCKSYYQLTX	-----RIPFIL	LTTAAS-----WIASECRVD	-----	LAGTIAI	GTATSGLLGCV-----
SLR0056 :	MSDIQTGQ--NOAKAROLLGKGAA	GESSIWKTRLO	WVPLWGVICAAASSGGYIW	-----SGPIMTGQTQ	-----	: 64
SLR1518 :	MTESSPLAP--STAPATRKWLWAAIK	-----PYTAY	AYPTVGSAAYGT	-----QWH-----GDUFTIF	-----SATIAIWNL	: 68
	M	P	G	1	L	: 71
SLR1736 :	NOMMDVDTRINKPNLPIANGDFS	AOGRIWV	GCG-----SLAIAWC	GLNGLITVGIS-----	IIGTAYSVPP	VLKRFSSLAAJC
SLR0926 :	NDLMDRDLPQVERTKORELLARA	RA-SMQVG	GVA-----LCAAGAFL	-----TPIPLS-----FW	TVRG	: 152
SLL1899 :	NCYDQDLYEMLTARPJPA	GKYSOPRHA	IPA-----ALEVN	-----TPIPLS-----FW	CVLANP	AYPGARVFVPVPO
SLR0056 :	NDFYDRDAINEPYKPSG	LISSPOTVQ	LL-----SIGVAY	-----FW	-----	: 150
SLR1518 :	NDYEDSDTGDIDRKAHSVNL	TSWRAQDW	-----V1	-----G	-----	: 71
	N	5D Did	G	1	6	: 156
SLR1736 :	VVNGLGLFTRIGLGYPP	ITIPINW	TAIAIFKDVPDM	-----Q1	-----	: 157
SLR0926 :	AMGEAVLIS	-----AVICD	TDATW	-----Q1	-----	: 157
SLL1899 :	AGSIPPVGA	-----VTGDS	WTGATVFM	-----Q1	-----	: 157
SLR0056 :	GASIAAPWNG-H	-----EGT	NPNTM	-----Q1	-----	: 157
SLR1518 :	LITEGP	-----AAAYYSOSQS	TMV	-----Q1	-----	: 157
	M	P	G	1	6	: 157
SLR1736 :	SLR1736 :	-----ASFYQRIWKLE	ESKTE	-----	-----	: 180
SLR0926 :	SLR0926 :	-----TEPKLYQ	-----IIGQ	-----	-----	: 180
SLL1899 :	SLL1899 :	-----QYIQLSAPT	-----IIGQ	-----	-----	: 180
SLR0056 :	SLR0056 :	-----LHQQLY	-----AATGGOF	-----VKAQWLKOAG	-----GDRDARG-L	: 180
SLR1518 :	SLR1518 :	-----YHQQYAT	-----LEND	-----KYO	-----ASACPF	: 180
	M	P	G	1	6	: 180
SLR1736 :	AMPLNTAII	SHICULLA	WVRSRDVHILESKTE	-----ASFYQRIWKLE	-----	: 241
SLR0926 :	MLNPN	YMSA	-----TPEKLYQ	-----IIGQ	-----	: 234
SLL1899 :	LHOLGLY	YMA	-----QYIQLSAPT	-----IIGQ	-----	: 241
SLR0056 :	YHQQYAT	-----L	-----LEND	-----KYO	-----ASACPF	: 263
SLR1518 :	QAPWQTLI	-----AS	-----POTFQDMYFLRN	-----IATCA	-----IGHAGI	: 246
	M	P	G	1	6	: 246
SLR1736 :	SLR1736 :	-----ASFYQRIWKLE	ESKTE	-----	-----	: 241
SLR0926 :	SLR0926 :	-----TEPKLYQ	-----IIGQ	-----	-----	: 234
SLL1899 :	SLL1899 :	-----QYIQLSAPT	-----IIGQ	-----	-----	: 241
SLR0056 :	SLR0056 :	-----LHQQLY	-----AATGGOF	-----VKAQWLKOAG	-----GDRDARG-L	: 263
SLR1518 :	SLR1518 :	-----YHQQYAT	-----LEND	-----KYO	-----ASACPF	: 246
	M	P	G	1	6	: 246
SLR1736 :	SLR1736 :	-----ASFYQRIWKLE	ESKTE	-----ASFYQRIWKLE	-----	: 308
SLR0926 :	SLR0926 :	-----TEPKLYQ	-----IIGQ	-----TEPKLYQ	-----	: 292
SLL1899 :	SLL1899 :	-----QYIQLSAPT	-----IIGQ	-----QYIQLSAPT	-----	: 316
SLR0056 :	SLR0056 :	-----LHQQLY	-----AATGGOF	-----VKAQWLKOAG	-----GDRDARG-L	: 324
SLR1518 :	SLR1518 :	-----YHQQYAT	-----LEND	-----KYO	-----ASACPF	: 307
	M	P	G	1	6	: 307

Figure 21

□□□□□□□□□□□□□□□□□□

ATPT2	:	*	20	*	40	*	60	*	80
SLR1736	:	-	-	-	-	-	-	-	-
ATPT3	:	-	-	-	-	-	-	-	-
SLR0926	:	-	-	-	-	-	-	-	-
ATPT4	:	-	-	-	-	-	-	-	-
SLL1899	:	-	-	-	-	-	-	-	-
ATPT12	:	-	-	-	-	-	-	-	-
SLR0056	:	-	-	-	-	-	-	-	-
ATPT8	:	-	-	-	-	-	-	-	-
SLR1518	:	-	-	-	-	-	-	-	-
		*	100	*	120	*	140	*	160
ATPT2	:	-	-	-	-	-	-	-	-
SLR1736	:	-	-	-	-	-	-	-	-
ATPT3	:	-	-	-	-	-	-	-	-
SLR0926	:	-	-	-	-	-	-	-	-
ATPT4	:	-	-	-	-	-	-	-	-
SLL1899	:	-	-	-	-	-	-	-	-
ATPT12	:	-	-	-	-	-	-	-	-
SLR0056	:	-	-	-	-	-	-	-	-
ATPT8	:	-	-	-	-	-	-	-	-
SLR1518	:	-	-	-	-	-	-	-	-
		*	200	*	220	*	240	*	260

ATPT2	:	*	AVIAALMINIYIVG	*	KVNKPYLPLASGEYSVNTGIA	*	VASEFS	*	MSFWLGIWIVGSWMPFLWALFISRFNIGTNTYS-INPILR
SLR1736	:	-	-	-	-	-	-	-	-
ATPT3	:	-	-	-	-	-	-	-	-
SLR0926	:	-	-	-	-	-	-	-	-
ATPT4	:	-	-	-	-	-	-	-	-
SLL1899	:	-	-	-	-	-	-	-	-
ATPT12	:	-	-	-	-	-	-	-	-
SLR0056	:	-	-	-	-	-	-	-	-
ATPT8	:	-	-	-	-	-	-	-	-
SLR1518	:	-	-	-	-	-	-	-	-
		*	200	*	220	*	240	*	260

6

Figure 22 1/2

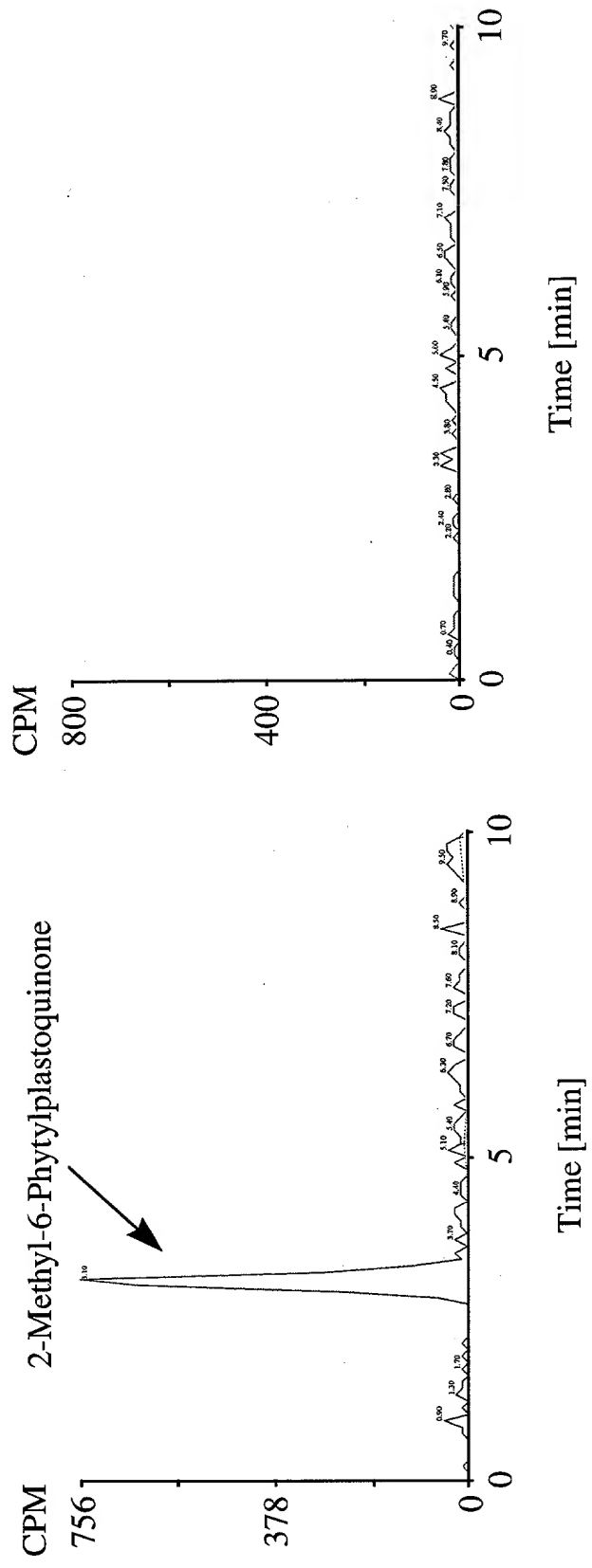
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□□□□□□□□□□□□□□□□□□

ATPT2	*	280	*	300	*	320	*	340	*	340	*	
SLR1736	:	WKR-FALVAAMCILLA	RAILYQIAFYTH-IOTHVFGRRPI	FTRPI	FATAFMSFESV	VIAI	FKDIDPDIIGDKIKI-FCTRSFSVTL	-QKR	:	313		
ATPT3	:	LKR-FSLJLAACILT	RGIVVNUGLFE-FRIGLGYPPT	ITPTW	-TFLFILVPTV	VATA	FKDVPDMGDRQ-FKTOQT	QIC-KQN	:	218		
SLR0926	:	SYP-LMKRFTFWMPQAFLGLT	TINNGAATG--	AT-	VYDTIYAHODKEDDVK-	VEYKXSTADRR-	DNT	--	:	328		
ATPT4	:	AVP-GAKRVFFVFPOL	LSIANGFAVIS	IS-	GFDTVYAMADREDDRR-	EVNNESSAEEFC-	-QYV	:	213			
SLL1899	:	VIT-PLKQLHPPINTW	GAVVGAIPPLG--	VA-	EPALALYFWOUPHEM	MAHALHLCRNDYAA-	GEYKMEIS	FDP--S	:	294		
ATPT12	:	VITHWLKRHTAQNTV	GGAAGSIPPING	VA-	EPALCO-SYNSTM	-EPALIFLW	LPHEFWALALMIKKDYAQ-	-DNVMPMLP	IAEKEKT	:	220	
SLR0056	:	IIS-APPLKLKONGW	GNFAAGASYISLWTAQAL	FECT	SWTPW	-LFALIFLW	TPHEFWALALMIKKDYAQ-	-DNVMPMLP	IAEKEKT	:	308	
ATPT8	:	IIS-APPLKLKONGW	GNYAAGASYTALEWNAH	FECT	ATDWT	-TFLYSLAG	GIAYVNDFKSVVEGR	RA-LOQIPAF-	-TET	:	242	
SLR1518	:	ELTSSTTBQRYSMDDYY	QKTYYKTASTISNSCKAVAVL	TGTTA	ATDWT	-TFLYSLAG	GIAYVNDFKSVVEGR	RO-LOQIPAF-	-IGT	:	231	
		TYQGPPFRRLGYLGELICLT	TFGPAT-AAYYSQSOSFESWNLLIT	PSVFGISTAI	DDLA-ACKRSPIVRL	-TKL	: 223					
ATPT2	*	360	*	380	*	400	*	420	*	440	*	
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SLR0926	:	KLWITGFGTGTASIGF	ALSGFSADLGWQYYASTAASGQ	GWO	GTADLSSGADC	--	-RKEVSNKWMFGAT	IFSGV	GRSFQ-	304		
ATPT4	:	GEAVGFFALTIGC	FYTGMIMLNPLYWLSTAAT	--	YGMV	OYIQLSAPTPEP-KIY	--	QGIEFGQV	IIIGFV	WLPNFS	:	
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SLR0056	:	AKWICGADMITQFLRNP	ASGGKPYIALVAL	--	IPQTFQKFLDPVKYDVK	--	YQASAQPFLVFGFLVTA	ASOH	--	379		
ATPT8	:	AAWICGADMIDVFOAQ	AGIAGLYTVHQQQLYATVLL	--	IPQTFQDMYFLRNPLEND	W	--YQASAQPFLVFGMLATG	AGHAGI-	:	303		
SLR1518	:	ITAPIFEAEEFPQ	REVDQVEKDPRNVDAILEY	GKSKG	CRARELAMEHANAAIGSLPTE	NDY	KRSRRALID	THRVTIRN	:	387		
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ATPT2	*											
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SLR0926	:	-----									-	
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ATPT8	:	K-----									-	
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Figure 22 2/2

□ □ □ □ □ □ □ □ □ □ □



*Synechocystis* 6803 wild type      *Synechocystis* slr1736 knockout

Figure 23

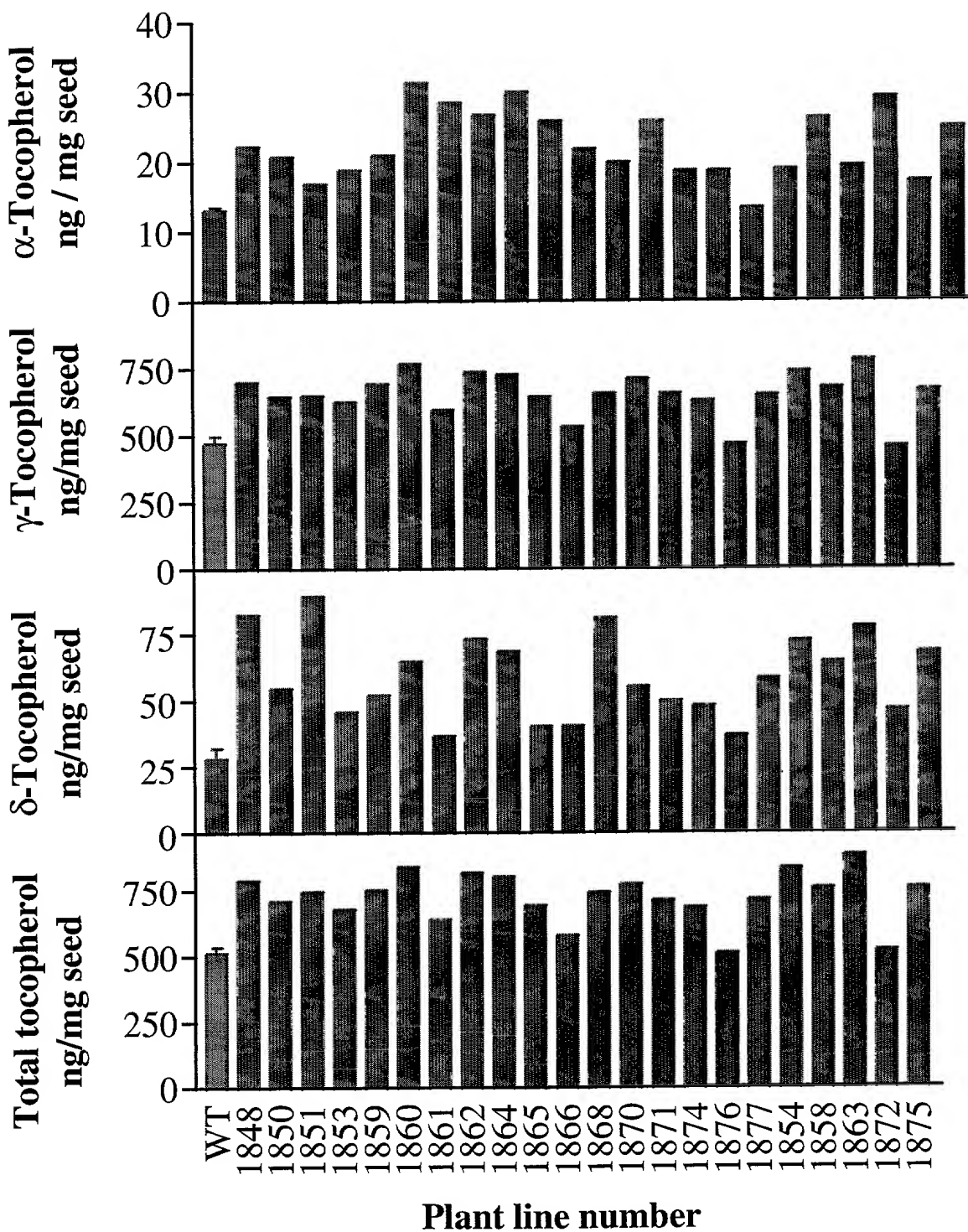


Figure 24

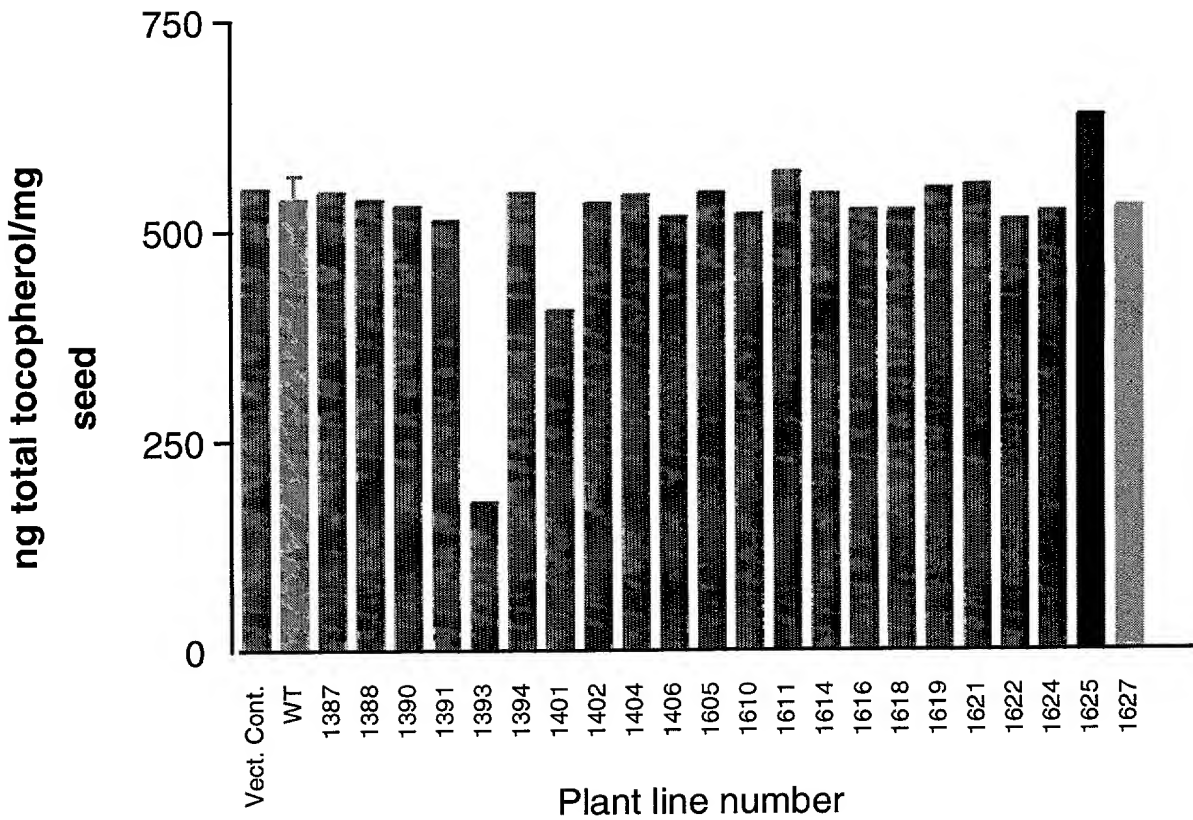


Figure 25

**SEQUENCE LISTING**

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85 85 90 95  
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Arg Tyr Ser Met Asp Tyr Tyr Met Gln Lys Thr Tyr Tyr Lys Thr Ala  
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Ser Leu Ile Ser Asn Ser Cys Lys Ala Val Ala Val Leu Thr Gly Gln  
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5 Thr Ala Glu Val Ala Val Leu Ala Phe Glu Tyr Gly Arg Asn Leu Gly  
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Leu Ala Phe Gln Leu Ile Asp Asp Ile Leu Asp Phe Thr Gly Thr Ser  
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Ala Ser Leu Gly Lys Gly Ser Leu Ser Asp Ile Arg His Gly Val Ile  
10 195 200 205  
Thr Ala Pro Ile Leu Phe Ala Met Glu Glu Phe Pro Gln Leu Arg Glu  
210 215 220  
Val Val Asp Gln Val Glu Lys Asp Pro Arg Asn Val Asp Ile Ala Leu  
225 230 235 240  
15 Glu Tyr Leu Gly Lys Ser Lys Gly Ile Gln Arg Ala Arg Glu Leu Ala  
245 250 255  
Met Glu His Ala Asn Leu Ala Ala Ala Ile Gly Ser Leu Pro Glu  
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Glu Ser Thr Asp Ile Val Thr Ser Glu Leu Arg Val Arg Gln Arg Gly  
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85 90 95  
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Ser Asp Ile Arg His Gly Val Ile Thr Ala Pro Ile Leu Phe Ala Met  
225 230 235 240  
15 Glu Glu Phe Pro Gln Leu Arg Glu Val Val Asp Gln Val Glu Lys Asp  
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Pro Arg Asn Val Asp Ile Ala Leu Glu Tyr Leu Gly Lys Ser Lys Gly  
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275 280 285  
20 Ala Ala Ile Gly Ser Leu Pro Glu Thr Asp Asn Glu Asp Val Lys Arg  
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15 ttgactgcga	caccaagctc	gacgagtaact	tactcaaaag	tttctacaag	acagcctctt	420
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35 cttgcgcggt	cgagatgatc	cacacaagct	ctctcattca	tgacgatctt	ccgtgcattgg	420
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35                    40                    45

Ser Pro Gly Arg Arg Phe Val Val Arg Ala Ala Glu Thr Asp Thr Asp  
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Lys Val Lys Ser Gln Thr Pro Asp Lys Ala Pro Ala Gly Gly Ser Ser  
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Ile Asn Gln Leu Leu Gly Ile Lys Gly Ala Ser Gln Glu Thr Asn Lys  
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Trp Lys Ile Arg Leu Gln Leu Thr Lys Pro Val Thr Trp Pro Pro Leu  
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10 Val Trp Gly Val Val Cys Gly Ala Ala Ala Ser Gly Asn Phe His Trp  
115                  120                  125

Thr Pro Glu Asp Val Ala Lys Ser Ile Leu Cys Met Met Met Ser Gly  
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Pro Cys Leu Thr Gly Tyr Thr Gln Thr Ile Asn Asp Trp Tyr Asp Arg  
15        145            150                  155                  160

Asp Ile Asp Ala Ile Asn Glu Pro Tyr Arg Pro Ile Pro Ser Gly Ala  
165                  170                  175

Ile Ser Glu Pro Glu Val Ile Thr Gln Val Trp Val Leu Leu Leu Gly  
180                  185                  190

Gly Leu Gly Ile Ala Gly Ile Leu Asp Val Trp Ala Gly His Thr Thr  
195                  200                  205

Pro Thr Val Phe Tyr Leu Ala Leu Gly Gly Ser Leu Leu Ser Tyr Ile  
210                  215                  220

Tyr Ser Ala Pro Pro Leu Lys Leu Lys Gln Asn Gly Trp Val Gly Asn  
225                  230                  235                  240

Phe Ala Leu Gly Ala Ser Tyr Ile Ser Leu Pro Trp Trp Ala Gly Gln  
245                  250                  255

Ala Leu Phe Gly Thr Leu Thr Pro Asp Val Val Val Leu Thr Leu Leu  
260                  265                  270

30 Tyr Ser Ile Ala Gly Leu Gly Ile Ala Ile Val Asn Asp Phe Lys Ser  
275                  280                  285

Val Glu Gly Asp Arg Ala Leu Gly Leu Gln Ser Leu Pro Val Ala Phe  
290                  295                  300

Gly Thr Glu Thr Ala Lys Trp Ile Cys Val Gly Ala Ile Asp Ile Thr  
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Gln Leu Ser Val Ala Gly Tyr Leu Leu Ala Ser Gly Lys Pro Tyr Tyr  
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Ala Leu Ala Leu Val Ala Leu Ile Ile Pro Gln Ile Val Phe Gln Phe  
340                  345                  350

40 Lys Tyr Phe Leu Lys Asp Pro Val Lys Tyr Asp Val Lys Tyr Gln Ala  
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840 tttgagatca ataatgatgc taaaatgaag agaacaagtc gcaggccact accctcagga  
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Val Ala Ser Phe Ser Ile Leu Ser Phe Trp Leu Gly Trp Val Val Gly  
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35 Ala Phe Phe Leu His Met Gln Thr His Val Tyr Lys Arg Pro Pro Val  
115 120 125  
Phe Ser Arg Pro Leu Ile Phe Ala Thr Ala Phe Met Ser Phe Phe Ser  
130 135 140  
Val Val Ile Ala Leu Phe Lys Asp Ile Pro Asp Ile Glu Gly Asp Lys  
40 145 150 155 160  
Val Phe Gly Ile Gln Ser Phe Ser Val Cys Leu Gly Gln Lys Pro Val  
165 170 175  
Phe Trp Thr Cys Val Thr Leu Leu Glu Ile Ala Tyr Gly Val Ala Leu

180 185 190  
Leu Val Gly Ala Ala Ser Pro Cys Leu Trp Ser Lys Ile Phe Thr Gly  
195 200 205  
Leu Gly His Ala Val Leu Ala Ser Ile Leu Trp Phe His Ala Lys Ser  
5 210 215 220  
Val Asp Leu Lys Ser Lys Ala Ser Ile Thr Ser Phe Tyr Met Phe Ile  
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atgtgggctaa caagtgttgg agttgcagga acagcttgcgtt tggcctggaa ggctaattggc 180  
ttggcagctg ggcttgcagc ttctaatctt gttctgtatg catttgcgtt aacggccgttg 240

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tgcttcggca gcttagcaact cagtggttac aatgctgacc ttggttggtg tttagtgtga 180  
tgcttgcgca aagaatggta tngttttac ttgatattga ctccagacct gaaatcatgt 240  
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30 agcgcacaac cattttt 257  
  
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40 aagaaggctt tttggatctg cggtggctt cttgagatgg cctacagcgt tgcgatactg 180  
atgggagcta ccttcctg tttgtggagc aaaacagcaa ccattcgctgg ccattccata 240  
cttgcgcga tcctatggag ctgcgcgcga tcgggtggact tgacgagcaa agccgcaata 300  
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368

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Phe Lys Asp Ile Pro Asp Ile Glu Gly Asp Arg Ile Phe Gly Ile Arg

20 25 30

Ser Phe Ser Val Arg Leu Gly Gln Lys Lys Val Phe Trp Ile Cys Val

35 40 45

15 Gly Leu Leu Glu Met Ala Tyr Ser Val Ala Ile Leu Met Gly Ala Thr

50 55 60

Ser Ser Cys Leu Trp Ser Lys Thr Ala Thr Ile Ala Gly His Ser Ile

65 70 75 80

Leu Ala Ala Ile Leu Trp Ser Cys Ala Arg Ser Val Asp Leu Thr Ser

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Lys Ala Ala Ile Thr Ser Phe Tyr Met Phe Ile Trp Lys Leu Phe Tyr

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Ala Glu Tyr Leu Leu Ile Pro Leu Val Arg

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gatatcattg tcttgactac tttgtacago atagctgggc tagggattgc tattgtaaat 180

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40 <212> PRT

<213> Synechocystis sp

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20 25 30  
5 Pro Ala Leu Trp Ala Val Cys Leu Ala Ala Gln Gly Leu Pro Pro Leu  
35 40 45  
Pro Leu Leu Gly Thr Ile Ala Leu Gly Thr Leu Ala Thr Ser Gly Leu  
50 55 60  
Gly Cys Val Val Asn Asp Leu Trp Asp Arg Asp Ile Asp Pro Gln Val  
10 65 70 75 80  
Glu Arg Thr Lys Gln Arg Pro Leu Ala Ala Arg Ala Leu Ser Val Gln  
85 90 95  
Val Gly Ile Gly Val Ala Leu Val Ala Leu Leu Cys Ala Ala Gly Leu  
100 105 110  
15 Ala Phe Tyr Leu Thr Pro Leu Ser Phe Trp Leu Cys Val Ala Ala Val  
115 120 125  
Pro Val Ile Val Ala Tyr Pro Gly Ala Lys Arg Val Phe Pro Val Pro  
130 135 140  
Gln Leu Val Leu Ser Ile Ala Trp Gly Phe Ala Val Leu Ile Ser Trp  
20 145 150 155 160  
Ser Ala Val Thr Gly Asp Leu Thr Asp Ala Thr Trp Val Leu Trp Gly  
165 170 175  
Ala Thr Val Phe Trp Thr Leu Gly Phe Asp Thr Val Tyr Ala Met Ala  
180 185 190  
25 Asp Arg Glu Asp Asp Arg Arg Ile Gly Val Asn Ser Ser Ala Leu Phe  
195 200 205  
Phe Gly Gln Tyr Val Gly Glu Ala Val Gly Ile Phe Phe Ala Leu Thr  
210 215 220  
Ile Gly Cys Leu Phe Tyr Leu Gly Met Ile Leu Met Leu Asn Pro Leu  
30 225 230 235 240  
Tyr Trp Leu Ser Leu Ala Ile Ala Ile Val Gly Trp Val Ile Gln Tyr  
245 250 255  
Ile Gln Leu Ser Ala Pro Thr Pro Glu Pro Lys Leu Tyr Gly Gln Ile  
260 265 270  
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Leu Gly Trp Leu  
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35 40 45  
Val Asp Leu Pro Lys Leu Leu Ile Thr Leu Leu Gly Gly Thr Leu Ala  
10 50 55 60  
Ala Ala Ser Ala Gln Thr Leu Asn Cys Ile Tyr Asp Gln Asp Ile Asp  
65 70 75 80  
Tyr Glu Met Leu Arg Thr Arg Ala Arg Pro Ile Pro Ala Gly Lys Val  
85 90 95  
15 Gln Pro Arg His Ala Leu Ile Phe Ala Leu Ala Leu Gly Val Leu Ser  
100 105 110  
Phe Ala Leu Leu Ala Thr Phe Val Asn Val Leu Ser Gly Cys Leu Ala  
115 120 125  
Leu Ser Gly Ile Val Phe Tyr Met Leu Val Tyr Thr His Trp Leu Lys  
130 135 140  
20 Arg His Thr Ala Gln Asn Ile Val Ile Gly Gly Ala Ala Gly Ser Ile  
145 150 155 160  
Pro Pro Leu Val Gly Trp Ala Ala Val Thr Gly Asp Leu Ser Trp Thr  
165 170 175  
25 Pro Trp Val Leu Phe Ala Leu Ile Phe Leu Trp Thr Pro Pro His Phe  
180 185 190  
Trp Ala Leu Ala Leu Met Ile Lys Asp Asp Tyr Ala Gln Val Asn Val  
195 200 205  
30 Pro Met Leu Pro Val Ile Ala Gly Glu Glu Lys Thr Val Ser Gln Ile  
210 215 220  
Trp Tyr Tyr Ser Leu Leu Val Val Pro Phe Ser Leu Leu Leu Val Tyr  
225 230 235 240  
Pro Leu His Gln Leu Gly Ile Leu Tyr Leu Ala Ile Ala Ile Ile Leu  
245 250 255  
35 Gly Gly Gln Phe Leu Val Lys Ala Trp Gln Leu Lys Gln Ala Pro Gly  
260 265 270  
Asp Arg Asp Leu Ala Arg Gly Leu Phe Lys Phe Ser Ile Phe Tyr Leu  
275 280 285  
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Gln Leu Val Ala Gln Met Gly Thr Leu Leu Leu Gly  
305 310 315

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Ile Arg Leu Gln Leu Met Lys Pro Ile Thr Trp Ile Pro Leu Ile Trp  
35 40 45  
Gly Val Val Cys Gly Ala Ala Ser Ser Gly Gly Tyr Ile Trp Ser Val  
50 55 60  
15 Glu Asp Phe Leu Lys Ala Leu Thr Cys Met Leu Leu Ser Gly Pro Leu  
65 70 75 80  
Met Thr Gly Tyr Thr Gln Thr Leu Asn Asp Phe Tyr Asp Arg Asp Ile  
85 90 95  
Asp Ala Ile Asn Glu Pro Tyr Arg Pro Ile Pro Ser Gly Ala Ile Ser  
100 105 110  
20 Val Pro Gln Val Val Thr Gln Ile Leu Ile Leu Leu Val Ala Gly Ile  
115 120 125  
Gly Val Ala Tyr Gly Leu Asp Val Trp Ala Gln His Asp Phe Pro Ile  
130 135 140  
25 Met Met Val Leu Thr Leu Gly Gly Ala Phe Val Ala Tyr Ile Tyr Ser  
145 150 155 160  
Ala Pro Pro Leu Lys Leu Lys Gln Asn Gly Trp Leu Gly Asn Tyr Ala  
165 170 175  
Leu Gly Ala Ser Tyr Ile Ala Leu Pro Trp Trp Ala Gly His Ala Leu  
30 180 185 190  
Phe Gly Thr Leu Asn Pro Thr Ile Met Val Leu Thr Leu Ile Tyr Ser  
195 200 205  
Leu Ala Gly Leu Gly Ile Ala Val Val Asn Asp Phe Lys Ser Val Glu  
210 215 220  
35 Gly Asp Arg Gln Leu Gly Leu Lys Ser Leu Pro Val Met Phe Gly Ile  
225 230 235 240  
Gly Thr Ala Ala Trp Ile Cys Val Ile Met Ile Asp Val Phe Gln Ala  
245 250 255  
Gly Ile Ala Gly Tyr Leu Ile Tyr Val His Gln Gln Leu Tyr Ala Thr  
40 260 265 270  
Ile Val Leu Leu Leu Leu Ile Pro Gln Ile Thr Phe Gln Asp Met Tyr  
275 280 285  
Phe Leu Arg Asn Pro Leu Glu Asn Asp Val Lys Tyr Gln Ala Ser Ala

290                    295                    300  
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20                    25                        30  
Val Pro Ile Thr Val Gly Ser Ala Val Ala Tyr Gly Leu Thr Gly Gln  
35                    40                        45  
Trp His Gly Asp Val Phe Thr Ile Phe Leu Leu Ser Ala Ile Ala Ile  
50                    55                        60  
Ile Ala Trp Ile Asn Leu Ser Asn Asp Val Phe Asp Ser Asp Thr Gly  
65                    70                        75                        80  
Ile Asp Val Arg Lys Ala His Ser Val Val Asn Leu Thr Gly Asn Arg  
85                    90                        95  
Asn Leu Val Phe Leu Ile Ser Asn Phe Phe Leu Leu Ala Gly Val Leu  
100                    105                      110  
Gly Leu Met Ser Met Ser Trp Arg Ala Gln Asp Trp Thr Val Leu Glu  
115                    120                      125  
Leu Ile Gly Val Ala Ile Phe Leu Gly Tyr Thr Tyr Gln Gly Pro Pro  
130                    135                      140  
Phe Arg Leu Gly Tyr Leu Gly Glu Leu Ile Cys Leu Ile Thr  
145                    150                      155                      160  
Phe Gly Pro Leu Ala Ile Ala Ala Tyr Tyr Ser Gln Ser Gln Ser  
165                    170                      175  
35 Phe Ser Trp Asn Leu Leu Thr Pro Ser Val Phe Val Gly Ile Ser Thr  
180                    185                      190  
Ala Ile Ile Leu Phe Cys Ser His Phe His Gln Val Glu Asp Asp Leu  
195                    200                      205  
Ala Ala Gly Lys Lys Ser Pro Ile Val Arg Leu Gly Thr Lys Leu Gly  
210                    215                      220  
Ser Gln Val Leu Thr Leu Ser Val Val Ser Leu Tyr Leu Ile Thr Ala  
225                    230                      235                      240  
Ile Gly Val Leu Cys His Gln Ala Pro Trp Gln Thr Leu Leu Ile Ile

245                    250                    255  
Ala Ser Leu Pro Trp Ala Val Gln Leu Ile Arg His Val Gly Gln Tyr  
              260                    265                    270  
His Asp Gln Pro Glu Gln Val Ser Asn Cys Lys Phe Ile Ala Val Asn  
5                275                    280                    285  
Leu His Phe Phe Ser Gly Met Leu Met Ala Ala Gly Tyr Gly Trp Ala  
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Gly Leu Gly  
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cctgcttccc tggatttagt gttcggcgct tggctggcct gcctgttggg taatgtgtac      180  
attgtcggcc tcaaccaatt gtggatgtg gacattgacc gcatcaataa gccgaatttg      240  
cccctagcta acggagattt ttctatcgcc cagggccgtt ggattgtggg actttgtggc      300  
gttgcttcct tggcgatcgc ctggggatta gggctatggc tggggctaac ggtgggcatt      360  
agtttgatta ttggcacggc ctattcggtg ccgcaggtaa ggttaaagcg cttttccctg      420  
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tttttttagaa ttgggtttagg ttatcccccc actttaataa ccccccattctg ggttttgact      540  
ttatattatct tagtttcac cgtggcgatc gccattttta aagatgtgcc agatatggaa      600  
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cggggAACCT taattttact cactgggtgt tattnagcca tggcaatctg gggcttatgg      720  
gcggctatgc cttaaatac tgctttctt attgttccc atttgtgctt attagcctta      780  
ctctggtggc ggagtgcaga tgtacactta gaaagcaaaa ccgaaattgc tagttttat      840  
cagtttattt ggaagctatt tttcttagag tacttgctgt atcccttggc tctgtggta      900  
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Ile Gly Thr Thr Leu Ser Val Trp Ala Val Tyr Leu Leu Thr Ile Leu  
              20                    25                    30

Gly Asp Gly Asn Ser Val Asn Ser Pro Ala Ser Leu Asp Leu Val Phe  
35 40 45  
Gly Ala Trp Leu Ala Cys Leu Leu Gly Asn Val Tyr Ile Val Gly Leu  
50 55 60  
5 Asn Gln Leu Trp Asp Val Asp Ile Asp Arg Ile Asn Lys Pro Asn Leu  
65 70 75 80  
Pro Leu Ala Asn Gly Asp Phe Ser Ile Ala Gln Gly Arg Trp Ile Val  
85 90 95  
Gly Leu Cys Gly Val Ala Ser Leu Ala Ile Ala Trp Gly Leu Gly Leu  
10 100 105 110  
Trp Leu Gly Leu Thr Val Gly Ile Ser Leu Ile Ile Gly Thr Ala Tyr  
115 120 125  
Ser Val Pro Pro Val Arg Leu Lys Arg Phe Ser Leu Leu Ala Ala Leu  
130 135 140  
15 Cys Ile Leu Thr Val Arg Gly Ile Val Val Asn Leu Gly Leu Phe Leu  
145 150 155 160  
Phe Phe Arg Ile Gly Leu Gly Tyr Pro Pro Thr Leu Ile Thr Pro Ile  
165 170 175  
Trp Val Leu Thr Leu Phe Ile Leu Val Phe Thr Val Ala Ile Ala Ile  
20 180 185 190  
Phe Lys Asp Val Pro Asp Met Glu Gly Asp Arg Gln Phe Lys Ile Gln  
195 200 205  
Thr Leu Thr Leu Gln Ile Gly Lys Gln Asn Val Phe Arg Gly Thr Leu  
210 215 220  
Ile Leu Leu Thr Gly Cys Tyr Leu Ala Met Ala Ile Trp Gly Leu Trp  
225 230 235 240  
Ala Ala Met Pro Leu Asn Thr Ala Phe Leu Ile Val Ser His Leu Cys  
245 250 255  
Leu Leu Ala Leu Leu Trp Trp Arg Ser Arg Asp Val His Leu Glu Ser  
30 260 265 270  
Lys Thr Glu Ile Ala Ser Phe Tyr Gln Phe Ile Trp Lys Leu Phe Phe  
275 280 285  
Leu Glu Tyr Leu Leu Tyr Pro Leu Ala Leu Trp Leu Pro Asn Phe Ser  
290 295 300  
35 Asn Thr Ile Phe  
305  
  
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40 <212> DNA  
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<400> 38

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	gaaaatcctg cttagcgatca tcattacggc ggccggctg tgcaaattt agggccggct	180
	acgaaaaaac aagaaaatca ggaagaccaa cttgttggc ggacattcc ctggtaaaa	240
5	aaatttggg ccagtccctcg ccagttgcc cttagggcatt gggggaaatg tagggataac	300
	aggcaggcga aaccctact ctccgaagaa tttttgccca cggtaagga aggttatcaa	360
	atccatcaa atcagcacca aggacaaatc attcatggcg atcgccattg tcgttggcag	420
	ttcacccgtag aaccggaagt aacttgggg agtcctaacc gatttcctcg ggctacagcg	480
	ggttggctt ccttttacc cttgtttagt cccgggttggc aaattcttt agcccaaggt	540
10	agagcgcacg gctggctgaa atggcagagg gaacagtatg aatttgacca cgccctagtt	600
	tatgccgaaa aaaattgggg tcactcctt ccctcccgct ggtttggct ccaagcaaat	660
	tatttcctg accatccagg actgagcgctc actgcccgtg gggggaaacg gattgttctt	720
	ggtcgccccg aagaggttagc ttaattggc ttacatcacc aaggttaattt ttacgaattt	780
	ggcccccggcc atggcacagt cacttggcaa gtagctccct gggccgttg gcaattaaaa	840
15	gcacagcaatg ataggtattt ggtcaagttt tccggaaaaa cagataaaaa aggcaatttt	900
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	ttgttatttgc aattgggatc tgggggtcac ggcctgatag tgcaagggga aacggacacc	1020
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25

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30	20	25	30	
	Ser Phe Ala Phe Met Tyr Ser Ile Glu Asn Pro Ala Ser Asp His His			
	35	40	45	
	Tyr Gly Gly Gly Ala Val Gln Ile Leu Gly Pro Ala Thr Lys Lys Gln			
	50	55	60	
35	Glu Asn Gln Glu Asp Gln Leu Val Trp Arg Thr Phe Pro Ser Val Lys			
	65	70	75	80
	Lys Phe Trp Ala Ser Pro Arg Gln Phe Ala Leu Gly His Trp Gly Lys			
	85	90	95	
40	Cys Arg Asp Asn Arg Gln Ala Lys Pro Leu Leu Ser Glu Glu Phe Phe			
	100	105	110	
	Ala Thr Val Lys Glu Gly Tyr Gln Ile His Gln Asn Gln His Gln Gly			
	115	120	125	
	Gln Ile Ile His Gly Asp Arg His Cys Arg Trp Gln Phe Thr Val Glu			

130                    135                    140  
Pro Glu Val Thr Trp Gly Ser Pro Asn Arg Phe Pro Arg Ala Thr Ala  
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Gly Trp Leu Ser Phe Leu Pro Leu Phe Asp Pro Gly Trp Gln Ile Leu  
5                      165                    170                    175  
Leu Ala Gln Gly Arg Ala His Gly Trp Leu Lys Trp Gln Arg Glu Gln  
180                    185                    190  
Tyr Glu Phe Asp His Ala Leu Val Tyr Ala Glu Lys Asn Trp Gly His  
195                    200                    205  
10 Ser Phe Pro Ser Arg Trp Phe Trp Leu Gln Ala Asn Tyr Phe Pro Asp  
210                    215                    220  
His Pro Gly Leu Ser Val Thr Ala Ala Gly Gly Glu Arg Ile Val Leu  
225                    230                    235                    240  
Gly Arg Pro Glu Glu Val Ala Leu Ile Gly Leu His His Gln Gly Asn  
15                      245                    250                    255  
Phe Tyr Glu Phe Gly Pro Gly His Gly Thr Val Thr Trp Gln Val Ala  
260                    265                    270  
Pro Trp Gly Arg Trp Gln Leu Lys Ala Ser Asn Asp Arg Tyr Trp Val  
275                    280                    285  
Lys Leu Ser Gly Lys Thr Asp Lys Lys Gly Ser Leu Val His Thr Pro  
290                    295                    300  
Thr Ala Gln Gly Leu Gln Leu Asn Cys Arg Asp Thr Thr Arg Gly Tyr  
305                    310                    315                    320  
Leu Tyr Leu Gln Leu Gly Ser Val Gly His Gly Leu Ile Val Gln Gly  
325                    330                    335  
Glu Thr Asp Thr Ala Gly Leu Glu Val Gly Gly Asp Trp Gly Leu Thr  
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